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(57) Abstract

The present invention is directed to 5' regulatory regions of an Arabidopsis oleosin gene. The 5' regulatory regions, when operably linked to either the coding sequence of a heterologous gene or a sequence complementary to a native plant gene, direct expression of the coding sequence or complementary sequence in a plant seed. The regulatory regions are useful in expression cassettes and expression vectors for the transformation of plants. Also provided are methods of modulating the levels of a heterologous gene such as a fatty acid synthesis or lipid metabolism gene by transforming a plant with the subject expression cassettes and expression vectors.

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AN OLEOSIN 5' REGULATORY REGION FOR THE MODIFICATION OF PLANT SEED LIPID COMPOSITION

BACKGROUND OF THE INVENTION

Seed oil content has traditionally been modified by plant breeding. The use of recombinant DNA technology to alter seed oil composition can accelerate this process and in some cases alter seed oils in a way that cannot be accomplished by breeding alone. The oil composition of *Brassica* has been significantly altered by modifying the expression of a number of lipid metabolism genes. Such manipulations of seed oil composition have focused on altering the proportion of endogenous component fatty acids. For example, antisense repression of the Δ12-desaturase gene in transgenic rapeseed has resulted in an increase in oleic acid of up to 83%. Topfer et al. 1995 *Science 268:681-686*.

There have been some successful attempts at modifying the composition of seed oil in transgenic plants by introducing new genes that allow the production of a fatty acid that the host plants were not previously capable of synthesizing. Van de Loo, et al. (1995 Proc. Natl. Acad. Sci USA 92:6743-6747) have been able to introduce a $\Delta 12$ -hydroxylase gene into transgenic tobacco, resulting in the introduction of a novel fatty acid, ricinoleic acid, into its seed oil. The reported accumulation was modest from plants carrying constructs in which transcription of the hydroxylase gene was under the control of the cauliflower mosaic virus (CaMV) 35S promoter.

Similarly, tobacco plants have been engineered to produce low levels of petroselinic acid by expression of an acyl-ACP desaturase from coriander (Cahoon et al. 1992 Proc. Natl. Acad. Sci USA 89:11184-11188).

The long chain fatty acids (C18 and larger), have significant economic value both as nutritionally and medically important foods and as industrial commodities (Ohlrogge, J.B. 1994 Plant Physiol. 104:821-826). Linoleic (18:2 Δ9,12) and α-linolenic acid (18:3 Δ9,12,15) are essential fatty acids found in many seed oils. The levels of these fatty-acids have been manipulated in oil seed crops through breeding and biotechnology (Ohlrogge, et al. 1991 Biochim. Biophys. Acta 1082:1-26; Topfer et al. 1995 Science 268:681-686). Additionally, the production of novel fatty acids in seed oils can be of considerable use in both human health and industrial applications.

Consumption of plant oils rich in γ linolenic acid (GLA) (18:3 Δ 6,9,12) is thought to
alleviate hypercholesterolemia and other related
clinical disorders which correlate with susceptibility
to coronary heart disease (Brenner R.R. 1976 Adv. Exp.
Med. Biol. 83:85-101). The therapeutic benefits of
dietary GLA may result from its role as a precursor to
prostaglandin synthesis (Weete, J.D. 1980 in Lipid
Biochemistry of Fungi and Other Organisms, eds. Plenum
Press, New York, pp. 59-62). Linoleic acid(18:2) (LA)
is transformed into gamma linolenic acid (18:3) (GLA)
by the enzyme Δ 6-desaturase.

Few seed oils contain GLA despite high contents of the precursor linoleic acid. This is due to the absence of A6-desaturase activity in most plants. For example, only borage (Borago officinalis), evening primrose (Oenothera biennis), and currants (Ribes nigrum) produce appreciable amounts of linolenic acid. Of these three species, only Oenothera and Borage are cultivated as a commercial source for GLA. It would be beneficial if agronomic seed oils could be engineered to produce GLA in significant quantities by introducing a heterologous Δ6-desaturase gene. It would also be beneficial if other expression products associated with fatty acid synthesis and lipid metabolism could be produced in plants at high enough levels so that commercial production of a particular expression product becomes feasible.

As disclosed in U.S. Patent No. 5,552,306, a cyanobacterial \$\Delta^6\$-desaturase gene has been recently isolated. Expression of this cyanobacterial gene in transgenic tobacco resulted in significant but low level GLA accumulation. (Reddy et al. 1996 Nature Biotech. 14:639-642). Applicant's copending U.S. Application Serial No. 08,366,779, discloses a \$\Delta^6\$-desaturase gene isolated from the plant Borago officinalis and its expression in tobacco under the control of the CaMV 35S promoter. Such expression resulted in significant but low level GLA and octadecatetraenoic acid (ODTA or OTA) accumulation in seeds. Thus, a need exists for a promoter which

functions in plants and which consistently directs high level expression of lipid metabolism genes in transgenic plant seeds.

Oleosins are abundant seed proteins associated with the phospholipid monolayer membrane of oil bodies. The first oleosin gene, L3, was cloned from maize by selecting clones whose in vitro translated products were recognized by an anti-L3 antibody (Vance et al. 1987 J. Biol. Chem. 262:11275-11279). Subsequently, different isoforms of oleosin genes from such different species as Brassica, soybean, carrot, pine, and Arabidopsis have been cloned (Huang, A.H.C., 1992, Ann. Reviews Plant Phys. and Plant Mol. Biol. 43:177-200; Kirik et al., 1996 Plant Mol. Biol. 31:413-417; Van Rooijen et al., 1992 Plant Mol. Biol. 18:1177-1179; Zou et al., Plant Mol. Biol. 31:429-433. Oleosin protein sequences predicted from these genes are highly conserved, especially for the central hydrophobic domain. All of these oleosins have the characteristic feature of three distinctive domains. An amphipathic domain of 40-60 amino acids is present at the N-terminus; a totally hydrophobic domain of 68-74 amino acids is located at the center; and an amphipathic α -helical domain of 33-40 amino acids is situated at the C-terminus (Huang, A.H.C. 1992).

The present invention provides 5' regulatory sequences from an oleosin gene which direct high level expression of lipid metabolism genes in transgenic plants. In accordance with the present invention,

chimeric constructs comprising an oleosin 5' regulatory region operably linked to coding sequence for a lipid metabolism gene such as a $\Delta 6$ -desaturase gene are provided. Transgenic plants comprising the subject chimeric constructs produce levels of GLA approaching the level found in those few plant species which naturally produce GLA such as evening primrose (Oenothera biennis).

SUMMARY OF THE INVENTION

The present invention is directed to 5' regulatory regions of an Arabidopsis oleosin gene. The 5' regulatory regions, when operably linked to either the coding sequence of a heterologous gene or sequence complementary to a native plant gene, direct expression of the heterologous gene or complementary sequence in a plant seed.

The present invention thus provides expression cassettes and expression vectors comprising an oleosin 5' regulatory region operably linked to a heterologous gene or a sequence complementary to a native plant gene.

Plant transformation vectors comprising the expression cassettes and expression vectors are also provided as are plant cells transformed by these vectors, and plants and their progeny containing the vectors.

In one embodiment of the invention, the heterologous gene or complementary gene sequence is a fatty acid synthesis gene or a lipid metabolism gene.

In another aspect of the present invention, a method is provided for producing a plant with increased levels of a product of a fatty acid synthesis or lipid metabolism gene.

In particular, there is provided a method for producing a plant with increased levels of a fatty acid synthesis or lipid metabolism gene by transforming a plant with the subject expression cassettes and expression vectors which comprise an oleosin 5' regulatory region and a coding sequence for a fatty acid synthesis or lipid metabolism gene.

In another aspect of the present invention, there is provided a method for cosuppressing a native fatty acid synthesis or lipid metabolism gene by transforming a plant with the subject expression cassettes and expression vectors which comprise an oleosin 5' regulatory region and a coding sequence for a fatty acid synthesis or lipid metabolism gene.

A further aspect of this invention provides a method of decreasing production of a native plant gene such as a fatty acid synthesis gene or a lipid metabolism gene by transforming a plant with an expression vector comprising a oleosin 5' regulatory region operably linked to a nucleic acid sequence complementary to a native plant gene.

Also provided are methods of modulating the levels of a heterologous gene such as a fatty acid synthesis or lipid metabolism gene by transforming a plant with the subject expression cassettes and expression vectors.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 depicts the nucleotide and corresponding amino acid sequence of the borage Δ6-desaturase gene (SEQ ID NO:1). The cytochrome b5 heme-binding motif is boxed and the putative metal binding, histidine rich motifs (HRMs) are underlined. The motifs recognized by the primers (PCR analysis) are underlined with dotted lines, i.e. tgg aaa tgg aac cat aa; and gag cat cat ttg ttt cc.

Fig. 2 is a dendrogram showing similarity of the borage $\Lambda 6$ -desaturase to other membrane-bound desaturases. The amino acid sequence of the borage $\Lambda 6$ -desaturase was compared to other known desaturases using Gene Works (IntelliGenetics). Numerical values correlate to relative phylogenetic distances between subgroups compared.

Fig. 3A provides a gas liquid chromatography profile of the fatty acid methyl esters (FAMES) derived from leaf tissue of a wild type tobacco 'Xanthi'.

Fig. 3B provides a gas liquid chromatography profile of the FAMES derived from leaf tissue of a tobacco plant transformed with the borage $\Delta 6$ -desaturase cDNA under transcriptional control of the CaMV 35S promoter (pAN2). Peaks corresponding to methyl linoleate (18:2), methyl γ -linolenate (18:3 γ), methyl α -linolenate (18:3 α), and methyl octadecatetraenoate (18:4) are indicated.

Fig. 4 is the nucleotide sequence and corresponding amino acid sequence of the oleosin AtS21 cDNA (SEO ID NO:3).

Fig. 5 is an acidic-base map of the predicted AtS21 protein generated by DNA Strider 1.2.

Fig. 6 is a Kyte-Doolittle plot of the predicted AtS21 protein generated by DNA Strider 1.2.

Fig. 7 is a sequence alignment of oleosins isolated from Arabidopsis. Oleosin sequences published or deposited in EMBL, BCM, NCBI databases were aligned to each other using GeneWorks® 2.3. Identical residues are boxed with rectangles. seven sequences fall into three groups. The first group includes AtS21 (SEQ ID NO:5), X91918 (SEQ ID NO:6) and Z29859 (SEQ ID NO:7). The second group includes X62352 (SEQ ID NO:8) and Ato13 (SEQ ID NO:9). The third group includes X91956 (SEQ ID NO:10) and L40954 (SEQ ID NO:11). Differences in amino acid residues within the same group are indicated by shadows. Ato2/Z54164 is identical to AtS21. Atol3 sequence (Accession No. Z541654 in EMBL database) is actually not disclosed in the EMBL database. 254165 Accession number designates the same sequence as Z54164 which is Atol2.

Fig. 8A is a Northern analysis of the AtS21 gene. An RNA gel blot containing ten micrograms of total RNA extracted from *Arabidopsis* flowers (F), leaves (L), roots (R), developing seeds (Se), and developing silique coats (Si) was hybridized with a probe made from the full-length AtS21 cDNA.

Fig. 8B is a Southern analysis of the AtS21 gene. A DNA gel blot containing ten micrograms of genomic DNA digested with BamHI (B), EcoRI (E), HindIII (H), SacI (S), and XbaI (X) was hybridized with a probe made from the full length AtS21 cDNA.

Fig. 9 is the nucleotide sequence of the SacI fragment of AtS21 genomic DNA (SEQ ID NO:12). The promoter and intron sequences are in uppercase. The fragments corresponding to AtS21 cDNA sequence are in lower case. The first ATG codon and a putative TATA box are shadowed. The sequence complementary to 21P primer for PCR amplification is boxed. A putative abscisic acid response element (ABRE) and two 14 bp repeats are underlined.

Fig. 10 is a map of AtS21 promoter/GUS
construct (pAN5).

Fig. 11A depicts AtS21/GUS gene expression in Arabidopsis bolt and leaves.

Fig. 11B depicts AtS21 GUS gene expression in Arabidopsis siliques.

Fig. 11C depicts AtS21 GUS gene expression in Arabidopsis developing seeds.

Figs. 11D through 11J depict AtS21 GUS gene expression in Arabidopsis developing embryos.

Fig. 11K depicts AtS21/GUS gene expression in Arabidopsis root and root hairs of a young seedling.

Fig. 11L depicts AtS21/GUS gene expression in Arabidopsis cotyledons and the shoot apex of a five day seedling.

Figs. 11M and 11N depict AtS21/GUS gene expression in *Arabidopsis* cotyledons and the shoot apex of 5-15 day seedlings.

Fig. 12A depicts AtS21/GUS gene expression in tobacco embryos and endosperm.

Fig. 12B depicts AtS21/GUS gene expression in germinating tobacco seeds.

Fig. 12C depicts AtS21/GUS gene expression in a 5 day old tobacco seedling.

Fig. 12D depicts AtS21/GUS gene expression in 5-15 day old tobacco seedlings.

Fig. 13A is a Northern analysis showing AtS21 mRNA levels in developing wild-type Arabidopsis seedlings. Lane 1 was loaded with RNA from developing seeds, lane 2 was loaded with RNA from seeds imbibed for 24-48 hours, lane 3: 3 day seedlings; lane 4: 4 day seedlings; lane 5: 5 day seedlings; lane 6: 6 day seedlings; lane 7; 9 day seedlings; lane 8: 12 day seedlings. Probe was labeled AtS21 cDNA. Exposure was for one hour at -80°C.

Fig. 13B is the same blot as Fig. 13A only exposure was for 24 hours at $-80\,^{\circ}\text{C}$.

Fig. 13C is the same blot depicted in Figs. 13A and 13B after stripping and hybridization with an Arabidopsis tubulin gene probe. The small band in each of lanes 1 and 2 is the remnant of the previous AtS21 probe. Exposure was for 48 hours at -80°C.

Fig. 14 is a graph comparing GUS activities expressed by the AtS21 and 35S promoters. GUS activities expressed by the AtS21 promoter in

developing Arabidopsis seeds and leaf are plotted side by side with those expressed by the 35S promoter. The GUS activities expressed by the AtS21 promoter in tobacco dry seed and leaf are plotted on the right side of the figure. GUS activity in tobacco leaf is so low that no column appears. "G-H" denotes globular to heart stage; "H-T" denotes heart to torpedo stage; "T-C" denotes torpedo to cotyledon stage; "Early C" denotes early cotyledon; "Late C" denotes late cotyledon. The standard deviations are listed in Table 2.

Fig. 15A is an RNA gel blot analysis carried out on 5 μg samples of RNA isolated from borage leaf, root, and 12 dpp embryo tissue, using labeled borage $\Delta 6$ -desaturase cDNA as a hybridization probe.

Fig. 15B depicts a graph corresponding to the Northern analysis results for the experiment shown in Fig. 15A.

Fig. 16A is a graph showing relative legumin RNA accumulation in developing borage embryos based on results of Northern blot.

Fig. 16B is a graph showing relative oleosin RNA accumulation in developing borage embryos based on results of Northern blot.

Fig. 16C is a graph showing relative $\Delta 6$ -desaturase RNA accumulation in developing borage embryos based on results of Northern blot.

Fig. 17 is a PCR analysis showing the presence of the borage delta 6-desaturase gene in transformed plants of oilseed rape. Lanes 1, 3 and 4

were loaded with PCR reactions performed with DNA from plants transformed with the borage delta 6-desaturase gene linked to the oleosin 5' regulatory region; lane 2: DNA from plant transformed with the borage delta 6-desaturase gene linked to the albumin 5' regulatory region; lanes 5 and 6: DNA from non-transformed plants; lane 7: molecular weight marker (1 kb ladder, Gibco BRL); lane 8: PCR without added template DNA; lane 9: control with DNA from Agrobacterium tumefaciens EHA 105 containing the plasmid pAN3 (i.e. the borage delta6-desaturase gene linked to the oleosin 5' regulatory region).

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides isolated nucleic acids encoding 5' regulatory regions from an Arabidopsis oleosin gene. In accordance with the present invention, the subject 5' regulatory regions, when operably linked to either a coding sequence of a heterologous gene or a sequence complementary to a native plant gene, direct expression of the coding sequence or complementary sequence in a plant seed. The oleosin 5' regulatory regions of the present invention are useful in the construction of an expression cassette which comprises in the 5' to 3' direction, a subject oleosin 5' regulatory region, a heterologous gene or sequence complementary to a native plant gene under control of the regulatory region and a 3' termination sequence. Such an expression cassette can be incorporated into a variety of autonomously replicating vectors in order to construct an expression vector.

It has been surprisingly found that plants transformed with the expression vectors of the present invention produce levels of GLA approaching the level found in those few plant species which naturally produce GLA such as evening primrose (Oenothera biennis).

As used herein, the term "cassette" refers to a nucleotide sequence capable of expressing a particular gene if said gene is inserted so as to be operably linked to one or more regulatory regions present in the nucleotide sequence. Thus, for example, the expression cassette may comprise a heterologous coding sequence which is desired to be expressed in a plant seed. The expression cassettes and expression vectors of the present invention are therefore useful for directing seed-specific expression of any number of heterologous genes. The term "seed-specific expression" as used herein, refers to expression in various portions of a plant seed such as the endosperm and embryo.

An isolated nucleic acid encoding a 5' regulatory region from an oleosin gene can be provided as follows. Oleosin recombinant genomic clones are isolated by screening a plant genomic DNA library with a cDNA (or a portion thereof) representing oleosin mRNA. A number of different oleosin cDNAs have been isolated. The methods used to isolate such cDNAs as well as the nucleotide and corresponding amino acid

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sequences have been published in Kirik et al. 1986

Plant Mol. Biol. 31:413-417; Zou et al. Plant Mol.

Biol. 31:429-433; Van Rooigen et al. 1992 Plant Mol.

Biol. 18:1177-1179.

Virtual subtraction screening of a tissue specific library using a random primed polymerase chain (RP-PCR) cDNA probe is another method of obtaining an oleosin cDNA useful for screening a plant genomic DNA library. Virtual subtraction screening refers to a method where a cDNA library is constructed from a target tissue and displayed at a low density so that individual cDNA clones can be easily separated. These cDNA clones are subtractively screened with driver quantities (i.e., concentrations of DNA to kinetically drive the hybridization reaction) of cDNA probes made from tissue or tissues other than the target tissue (i.e. driver tissue). The hybridized plaques represent genes that are expressed in both the target and the driver tissues; the unhybridized plaques represent genes that may be target tissuespecific or low abundant genes that can not be detected by the driver cDNA probe. The unhybridized cDNAs are selected as putative target tissue-specific genes and further analyzed by one-pass sequencing and Northern hybridization.

Random primed PCR (RP-PCR) involves synthesis of large quantities of cDNA probes from a trace amount of cDNA template. The method combines the amplification power of PCR with the representation of random priming to simultaneously amplify and label double-stranded cDNA in a single tube reaction.

Methods considered useful in obtaining oleosin genomic recombinant DNA are provided in Sambrook et al. 1989, in Molecular Cloning: A Laboratory Manual, Cold Spring Harbor, NY, for example, or any of the myriad of laboratory manuals on recombinant DNA technology that are widely available. To determine nucleotide sequences, a multitude of techniques are available and known to the ordinarily skilled artisan. For example, restriction fragments containing an oleosin regulatory region can be subcloned into the polylinker site of a sequencing vector such as pBluescript (Stratagene). These pBluescript subclones can then be sequenced by the double-stranded dideoxy method (Chen and Seeburg, 1985, DNA 4:165).

In a preferred embodiment, the oleosin regulatory region comprises nucleotides 1-1267 of Fig. 9 (SEQ ID NO:12). Modifications to the oleosin regulatory region as set forth in SEQ ID NO:12 which maintain the characteristic property of directing seed-specific expression, are within the scope of the present invention. Such modifications include insertions, deletions and substitutions of one or more nucleotides.

The 5' regulatory region of the present invention can be derived from restriction endonuclease or exonuclease digestion of an oleosin genomic clone. Thus, for example, the known nucleotide or amino acid

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sequence of the coding region of an isolated oleosin gene (e.g. Fig. 7) is aligned to the nucleic acid or deduced amino acid sequence of an isolated oleosin genomic clone and 5' flanking sequence (i.e., sequence upstream from the translational start codon of the coding region) of the isolated oleosin genomic clone located.

The oleosin 5' regulatory region as set forth in SEQ ID NO:12 (nucleotides 1-1267 of Fig. 9) may be generated from a genomic clone having either or both excess 5' flanking sequence or coding sequence by exonuclease III-mediated deletion. This is accomplished by digesting appropriately prepared DNA with exonuclease III (exoIII) and removing aliquots at increasing intervals of time during the digestion. The resulting successively smaller fragments of DNA may be sequenced to determine the exact endpoint of the deletions. There are several commercially available systems which use exonuclease III (exoIII) to create such a deletion series, e.g. Promega Biotech, "Erase-A-Base" system. Alternatively, PCR primers can be defined to allow direct amplification of the subject 5' regulatory regions.

Using the same methodologies, the ordinarily skilled artisan can generate one or more deletion fragments of nucleotides 1-1267 as set forth in SEQ ID NO:12. Any and all deletion fragments which comprise a contiguous portion of nucleotides set forth in SEQ ID NO:12 and which retain the capacity to

direct seed-specific expression are contemplated by the present invention.

The identification of oleosin 5' regulatory sequences which direct seed-specific expression comprising nucleotides 1-1267 of SEQ ID NO:12 and modifications or deletion fragments thereof, can be accomplished by transcriptional fusions of specific sequences with the coding sequences of a heterologous gene, transfer of the chimeric gene into an appropriate host, and detection of the expression of the heterologous gene. The assay used to detect expression depends upon the nature of the heterologous sequence. For example, reporter genes, exemplified by chloramphenicol acetyl transferase and \$-glucuronidase (GUS), are commonly used to assess transcriptional and translational competence of chimeric constructions. Standard assays are available to sensitively detect the reporter enzyme in a transgenic organism. The β glucuronidase (GUS) gene is useful as a reporter of promoter activity in transgenic plants because of the high stability of the enzyme in plant cells, the lack of intrinsic β-glucuronidase activity in higher plants and availability of a quantitative fluorimetric assay and a histochemical localization technique. Jefferson et al. (1987 EMBO J 6:3901) have established standard procedures for biochemical and histochemical detection of GUS activity in plant tissues. Biochemical assays are performed by mixing plant tissue lysates with 4methylumbelliferyl- β -D-glucuronide, a fluorimetric substrate for GUS, incubating one hour at 37°C, and

then measuring the fluorescence of the resulting 4-methyl-umbelliferone. Histochemical localization for GUS activity is determined by incubating plant tissue samples in 5-bromo-4-chloro-3-indolyl-glucuronide (X-Gluc) for about 18 hours at 37°C and observing the staining pattern of X-Gluc. The construction of such chimeric genes allows definition of specific regulatory sequences and demonstrates that these sequences can direct expression of heterologous genes in a seed-specific manner.

Another aspect of the invention is directed to expression cassettes and expression vectors (also termed herein "chimeric genes") comprising a 5' regulatory region from an oleosin gene which directs seed specific expression operably linked to the coding sequence of a heterologous gene such that the regulatory element is capable of controlling expression of the product encoded by the heterologous gene. The heterologous gene can be any gene other than oleosin. If necessary, additional regulatory elements or parts of these elements sufficient to cause expression resulting in production of an effective amount of the polypeptide encoded by the heterologous gene are included in the chimeric constructs.

Accordingly, the present invention provides chimeric genes comprising sequences of the oleosin 5' regulatory region that confer seed-specific expression which are operably linked to a sequence encoding a heterologous gene such as a lipid metabolism enzyme.

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Examples of lipid metabolism genes useful for practicing the present invention include lipid desaturases such as $\Delta 6$ -desaturases, $\Delta 12$ -desaturases, $\Delta 15$ -desaturases and other related desaturases such as stearoyl-ACP desaturases, acyl carrier proteins (ACPs), thioesterases, acetyl transacylases, acetyl-coA carboxylases, ketoacyl-synthases, malonyl transacylases, and elongases. Such lipid metabolism genes have been isolated and characterized from a number of different bacteria and plant species. Their nucleotide coding sequences as well as methods of isolating such coding sequences are disclosed in the published literature and are widely available to those of skill in the art.

In particular, the A6-desaturase genes disclosed in U.S. Patent No. 5,552,306 and applicants' copending U.S. Application Serial No. 08/366,779 filed December 30, 1994 and incorporated herein by reference, are contemplated as lipid metabolism genes particularly useful in the practice of the present invention.

The chimeric genes of the present invention are constructed by ligating a 5' regulatory region of a oleosin genomic DNA to the coding sequence of a heterologous gene. The juxtaposition of these sequences can be accomplished in a variety of ways. In a preferred embodiment the order of the sequences, from 5' to 3', is an oleosin 5' regulatory region (including a promoter), a coding sequence, and a

termination sequence which includes a polyadenylation site.

Standard techniques for construction of such chimeric genes are well known to those of ordinary skill in the art and can be found in references such as Sambrook et al.(1989). A variety of strategies are available for ligating fragments of DNA, the choice of which depends on the nature of the termini of the DNA fragments. One of ordinary skill in the art recognizes that in order for the heterologous gene to be expressed, the construction requires promoter elements and signals for efficient polyadenylation of the transcript. Accordingly, the oleosin 5' regulatory region that contains the consensus promoter sequence known as the TATA box can be ligated directly to a promoterless heterologous coding sequence.

The restriction or deletion fragments that contain the oleosin TATA box are ligated in a forward orientation to a promoterless heterologous gene such as the coding sequence of β -glucuronidase (GUS). The skilled artisan will recognize that the subject oleosin 5' regulatory regions can be provided by other means, for example chemical or enzymatic synthesis. The 3' end of a heterologous coding sequence is optionally ligated to a termination sequence comprising a polyadenylation site, exemplified by, but not limited to, the nopaline synthase polyadenylation site, or the octopine T-DNA gene 7 polyadenylation site. Alternatively, the polyadenylation site can be provided by the heterologous gene.

The present invention also provides methods of increasing levels of heterologous genes in plant seeds. In accordance with such methods, the subject expression cassettes and expression vectors are introduced into a plant in order to effect expression of a heterologous gene. For example, a method of producing a plant with increased levels of a product of a fatty acid synthesis or lipid metabolism gene is provided by transforming a plant cell with an expression vector comprising an oleosin 5' regulatory region operably linked to a fatty acid synthesis or lipid metabolism gene and regenerating a plant with increased levels of the product of said fatty acid synthesis or lipid metabolism gene.

Another aspect of the present invention provides methods of reducing levels of a product of a gene which is native to a plant which comprises transforming a plant cell with an expression vector comprising a subject oleosin regulatory region operably linked to a nucleic acid sequence which is complementary to the native plant gene. In this manner, levels of endogenous product of the native plant gene are reduced through the mechanism known as antisense regulation. Thus, for example, levels of a product of a fatty acid synthesis gene or lipid metabolism gene are reduced by transforming a plant with an expression vector comprising a subject oleosin 5' regulatory region operably linked to a nucleic acid sequence which is complementary to a nucleic acid

sequence coding for a native fatty acid synthesis or lipid metabolism gene.

The present invention also provides a method of cosuppressing a gene which is native to a plant which comprises transforming a plant cell with an expression vector comprising a subject oleosin 5' regulatory region operably linked to a nucleic acid sequence coding for the native plant gene. manner, levels of endogenous product of the native plant gene are reduced through the mechanism known as cosuppression. Thus, for example, levels of a product of a fatty acid synthesis gene or lipid metabolism gene are reduced by transforming a plant with an expression vector comprising a subject oleosin 5' regulatory region operably linked to a nucleic acid sequence coding for a native fatty acid synthesis or lipid metabolism gene native to the plant. Although the exact mechanism of cosuppression is not completely understood, one skilled in the art is familiar with published works reporting the experimental conditions and results associated with cosuppression (Napoli et al. 1990 The Plant Cell 2:270-289; Van der Krol 1990 The Plant Cell 2:291-299.

To provide regulated expression of the heterologous or native genes, plants are transformed with the chimeric gene constructions of the invention. Methods of gene transfer are well known in the art. The chimeric genes can be introduced into plants by leaf disk transformation-regeneration procedure as described by Horsch et al. 1985 Science 227:1229.

Other methods of transformation such as protoplast culture (Horsch et al. 1984 Science 223:496, DeBlock et al. 1984 EMBO J. 2:2143, Barton et al. 1983, Cell 32:1033) can also be used and are within the scope of this invention. In a preferred embodiment, plants are transformed with Agrobacterium-derived vectors such as those described in Klett et al. (1987) Annu. Rev. Plant Physiol. 38:467. Other well-known methods are available to insert the chimeric genes of the present invention into plant cells. Such alternative methods include biolistic approaches (Klein et al. 1987 Nature 327:70), electroporation, chemically-induced DNA uptake, and use of viruses or pollen as vectors.

When necessary for the transformation method, the chimeric genes of the present invention can be inserted into a plant transformation vector, e.g. the binary vector described by Bevan, M. 1984 Nucleic Acids Res. 12:8711-8721. Plant transformation vectors can be derived by modifying the natural gene transfer system of Agrobacterium tumefaciens. natural system comprises large Ti (tumor-inducing) plasmids containing a large segment, known as T-DNA, which is transferred to transformed plants. Another segment of the Ti plasmid, the vir region, is responsible for T-DNA transfer. The T-DNA region is bordered by terminal repeats. In the modified binary vectors, the tumor inducing genes have been deleted and the functions of the vir region are utilized to transfer foreign DNA bordered by the T-DNA border sequences. The T-region also contains a selectable

marker for antibiotic resistance, and a multiple cloning site for inserting sequences for transfer. Such engineered strains are known as "disarmed" A. tumefaciens strains, and allow the efficient transfer of sequences bordered by the T-region into the nuclear genome of plants.

Surface-sterilized leaf disks and other susceptible tissues are inoculated with the "disarmed" foreign DNA-containing A. tumefaciens, cultured for a number of days, and then transferred to antibiotic-containing medium. Transformed shoots are then selected after rooting in medium containing the appropriate antibiotic, and transferred to soil. Transgenic plants are pollinated and seeds from these plants are collected and grown on antibiotic medium.

Expression of a heterologous or reporter gene in developing seeds, young seedlings and mature plants can be monitored by immunological, histochemical or activity assays. As discussed herein, the choice of an assay for expression of the chimeric gene depends upon the nature of the heterologous coding region. For example, Northern analysis can be used to assess transcription if appropriate nucleotide probes are available. If antibodies to the polypeptide encoded by the heterologous gene are available, Western analysis and immunohistochemical localization can be used to assess the production and localization of the polypeptide. Depending upon the heterologous gene, appropriate biochemical assays can be used. For example,

acetyltransferases are detected by measuring acetylation of a standard substrate. The expression of a lipid desaturase gene can be assayed by analysis of fatty acid methyl esters (FAMES).

Another aspect of the present invention provides transgenic plants or progeny of these plants containing the chimeric genes of the invention. monocotyledonous and dicotyledonous plants are contemplated. Plant cells are transformed with the chimeric genes by any of the plant transformation methods described above. The transformed plant cell, usually in the form of a callus culture, leaf disk, explant or whole plant (via the vacuum infiltration method of Bechtold et al. 1993 C.R. Acad. Sci. Paris, 316:1194-1199) is regenerated into a complete transgenic plant by methods well-known to one of ordinary skill in the art (e.g. Horsch et al. 1985 Science 227:1129). In a preferred embodiment, the transgenic plant is sunflower, cotton, oil seed rape, maize, tobacco, Arabidopsis, peanut or soybean. progeny of transformed plants inherit the chimeric genes, seeds or cuttings from transformed plants are used to maintain the transgenic line.

The following examples further illustrate the invention.

EXAMPLE 1

Isolation of Membrane-Bound Polysomal RNA and Construction of Borage cDNA Library

Membrane-bound polysomes were isolated from borage seeds 12 days post pollination (12 DPP)using the protocol established for peas by Larkins and Davies (1975 Plant Phys. 55: 749-756). RNA was extracted from the polysomes as described by Mechler (1987 Methods in Enzymology 152: 241-248, Academic Press). Poly-A* RNA was isolated from the membrane bound polysomal RNA using Oligotex-dTTM beads (Qiagen).

Corresponding cDNA was made using Stratagene's ZAP cDNA synthesis kit. The cDNA library was constructed in the lambda ZAP II vector (Stratagene) using the lambda ZAP II kit. The primary library was packaged with Gigapack II Gold packaging extract (Stratagene).

EXAMPLE 2

Isolation of a A-6 Desaturase cDNA from Borage

Hybridization protocol

The amplified borage cDNA library was plated at low density (500 pfu on 150 mm petri dishes). Highly prevalent seed storage protein cDNAs were reduced (subtracted from the total cDNAs) by screening with the corresponding cDNAs.

Hybridization probes for screening the borage cDNA library were generated by using random primed DNA synthesis as described by Ausubel et al (1994 Current Protocols in Molecular Biology, Wiley Interscience, N.Y.) and corresponded to previously identified abundantly expressed seed storage protein cDNAs. Unincorporated nucleotides were removed by use of a G-50 spin column (Boehringer Manheim). denatured for hybridization by boiling in a water bath for 5 minutes, then quickly cooled on ice. Nitrocellulose filters carrying fixed recombinant bacteriophage were prehybridized at 60°C for 2-4 hours in hybridization solution [4X SET (600 mM NaCl, 80 mM Tris-HCl, 4 mM Na, EDTA; pH 7.8), 5X Denhardt's reagent (0.1% bovine serum albumin, 0.1% Ficoll, and 0.1% polyvinylpyrolidone), 100 µg/ml denatured salmon sperm DNA, 50 µg/ml polyadenine and 10 ug/ml polycytidine]. This was replaced with fresh hybridization solution to which denatured radioactive probe (2 ng/ml hybridization solution) was added. The filters were incubated at 60°C with agitation overnight. Filters

were washed sequentially in 4X, 2X, and 1X SET (150 mM NaCl, 20 mM Tris-HCl, 1 mM Na₂EDTA; pH7.8) for 15 minutes each at 60°C. Filters were air dried and then exposed to X-ray film for 24 hours with intensifying screens at -80°C.

Non-hybridizing plaques were excised using Stratagene's excision protocol and reagents.
Resulting bacterial colonies were used to inoculate liquid cultures and were either sequenced manually or by an ABI automated sequencer.

Random Sequencing of cDNAs from a Borage Seed 12 (DPP) Membrane-Bound Polysomal Library

Each cDNA corresponding to a non-hybridizing plaque was sequenced once and a sequence tag generated from 200-300 base pairs. All sequencing was performed by cycle sequencing (Epicentre). Over 300 expressed sequence tags (ESTs) were generated. Each sequence tag was compared to the GenBank database using the BLAST algorithm (Altschul et al. 1990 J. Mol. Biol. 215:403-410). A number of lipid metabolism genes, including the Δ6-desaturase were identified.

Database searches with the cDNA clone designated mbp-65 using BLASTX with the GenBank database resulted in a significant match to the previously isolated *Synechocystis* $\Delta 6$ -desaturase. It was determined however, that mbp-65 was not a full length cDNA. A full length cDNA was isolated using mbp-65 to screen the borage membrane-bound polysomal library. The resultant clone was designated pAN1 and the cDNA insert of pAN1 was sequenced by the cycle

sequencing method. The amino acid sequence deduced from the open reading frame (Fig. 1, SEQ ID NO:1) was compared to other known desaturases using Geneworks (IntelligGenetics) protein alignment program. This alignment indicated that the cDNA insert of pAN1 was the borage $\Delta 6$ -desaturase gene.

The resulting dendrogram (Figure 2) shows that $\Delta^{15}\text{-desaturases}$ and $\Delta^{12}\text{-desaturases}$ comprise two groups. The newly isolated borage sequence and the previously isolated Synechocystis Δ^6 -desaturase (U.S. Patent No. 5,552,306) formed a third distinct group. A comparison of amino acid motifs common to desaturases and thought to be involved catalytically in metal binding illustrates the overall similarity of the protein encoded by the borage gene to desaturases in general and the Synechocystis Δ^6 -desaturase in particular (Table 1). At the same time, comparison of the motifs in Table 1 indicates definite differences between this protein and other plant desaturases. Furthermore, the borage sequence is also distinguished from known plant membrane associated fatty acid desaturases by the presence of a heme binding motif conserved in cytochrome b, proteins (Schmidt et al. 1994 Plant Mol. Biol. 26:631-642) (Figure 1). while these results clearly suggested that the isolated cDNA was a borage Δ^6 -desaturase gene, further confirmation was necessary. To confirm the identity of the borage $\Delta 6$ -desaturase cDNA, the cDNA insert from pAN1 was cloned into an expression cassette for stable expression. The vector pBI121 (Jefferson et al. 1987

EMBO J. <u>6</u>:3901-3907) was prepared for ligation by digestion with BamHI and EcoICR I (an isoschizomer of SacI which leaves blunt ends; available from Promega) which excises the GUS coding region leaving the 35S promoter and NOS terminator intact. The borage Δ^6 -desaturase cDNA was excised from the recombinant plasmid (pAN1) by digestion with BamHI and XhoI. The XhoI end was made blunt by performing a fill-in reaction catalyzed by the Klenow fragment of DNA polymerase T. This fragment was then cloned into the BamHI/EcoICR I sites of pBI121.1, resulting in the plasmid pAN2.

TABLE 1

COMPARISON OF COMMON AMINO ACID MOTIFS IN MEMBRANE-BOUND DESATURASES

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- 4	NO:21)	HNYLHH (SEQ. ID. NO:22)	NO:23)	ID. NO:23)	NO:23)	NO:23)	NO:23)	NO:24)	NO:24)	NO:25)	NO:25)	NO:26)	NO:27)	NO:28)
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Desaturase	Borage o	Sympocyetis o'	Arab, chloroplast 513	Rice 013	Glycine chloroplast A:3	Arab. fad3 (013)	Brassica fad 3 (413	Borage A ¹² (P1-81)*	arab. fad2 (Δ^{13})	Arab, chloroplast A11	Giveine plastid A ¹¹	Spinach plastidial n-6	Symposition 13	Anabaena O ¹³

*P1-81 is a full length cDNA which was identified by EST analysis and shows high similarity to the Arbidopsis A12 desaturase (fad2)

EXAMPLE 3

Production of Transgenic Plants and Preparation and Analysis of Fatty Acid Methyl Esters (FAMEs)

The expression plasmid, pAN2 was used to transform tobacco (Nicotiana tabacum cv. xanthi) via Agrobacterium tumefaciens according to standard procedures (Horsch, et al. 1985 Science 227:1229-1231; Bogue et al. 1990 Mol. Gen. Genet. 221:49-57) except that the initial transformants were selected on 100 μ g/ml kanamycin.

Tissue from transgenic plants was frozen in liquid nitrogen and lyophilized overnight. FAMEs were prepared as described by Dahmer, et al. (1989) J. Amer. Oil. Chem. Soc. 66: 543-548. In some cases, the solvent was evaporated again, and the FAMEs were resuspended in ethyl acetate and extracted once with deionized water to remove any water soluble contaminants. FAMEs were analyzed using a Tracor-560 gas liquid chromatograph as previously described (Reddy et al. 1996 Nature Biotech. 14:639-642).

As shown in Figure. 3, transgenic tobacco leaves containing the borage cDNA produced both GLA and octadecatetraenoic acid (OTA) (18:4 $\Delta6.9.12.15$). These results thus demonstrate that the isolated cDNA encodes a borage $\Delta6$ -desaturase.

PCT/US98/07179

EXAMPLE 4

Expression of \(\Delta 6 \)-desaturase in Borage

The native expression of A6-desaturase was examined by Northern Analysis of RNA derived from borage tissues. RNA was isolated from developing borage embryos following the method of Chang et al. 1993 Plant Mol. Biol. Rep. 11:113-116. RNA was electrophoretically separated on formaldehyde-agarose gels, blotted to nylon membranes by capillary transfer, and immobilized by baking at 80°C for 30 minutes following standard protocols (Brown T., 1996 in Current Protocols in Molecular Biology, eds. Auselbel, et al. [Greene Publishing and Wiley-Interscience, New York] pp. 4.9.1-4.9.14.). filters were preincubated at 42°C in a solution containing 50% deionized formamide, 5X Denhardt's reagent, 5X SSPE (900 mM NaCl; 50mM Sodium phosphate, pH7.7; and 5 mM EDTA), 0.1% SDS, and 200 µg/ml denatured salmon sperm DNA. After two hours, the filters were added to a fresh solution of the same composition with the addition of denatured radioactive hybridization probe. In this instance, the probes used were borage legumin cDNA (Fig. 16A), borage oleosin cDNA (Fig. 16B), and borage Δ6-desaturase cDNA (pAN1, Example 2) (Fig. 16C). The borage legumin and oleosin cDNAs were isolated by EST cloning and identified by comparison to the GenBank database using the BLAST algorithm as described in Example 2. Loading variation was corrected by normalizing to

levels of borage EF1 α mRNA. EF1 α mRNA was identified by correlating to the corresponding cDNA obtained by the EST analysis described in Example 2. The filters were hybridized at 42°C for 12-20 hours, then washed as described above (except that the temperature was 65°C), air dried, and exposed to X-ray film.

As depicted in Figs. 15A and 15B, Δ6-desaturase is expressed primarily in borage seed. Borage seeds reach maturation between 18-20 days post pollination (dpp). Δ6-desaturase mRNA expression occurs throughout the time points collected (8-20 dpp), but appears maximal from 10-16 days post pollination. This expression profile is similar to that seen for borage oleosin and 12S seed storage protein mRNAs (Figs. 16A, 16B, and 16C).

EXAMPLE 5

Isolation and Characterization of a Novel Oleosin cDNA

The oleosin cDNA (AtS21) was isolated by virtual subtraction screening of an Arabidopsis developing seed cDNA library using a random primed polymerase chain reaction (RP-PCR) cDNA probe derived from root tissue.

RNA PREPARATION

Arabidopsis thaliana Landsberg erecta plants were grown under continuous illumination in a vermiculite/soil mixture at ambient temperature (22°C). Siliques 2-5 days after flowering were dissected to separately collect developing seeds and silique coats. Inflorescences containing initial flower buds and fully opened flowers, leaves, and whole siliques one or three days after flowering were also collected. Roots were obtained from seedlings that had been grown in Gamborg B, liquid medium (GIBCO BRL) for two weeks. The seeds for root culture were previously sterilized with 50% bleach for five minutes and rinsed with water extensively. All tissues were frozen in liquid nitrogen and stored at -80°C until use. Total RNAs were isolated following a hot phenol/SDS extraction and LiCl precipitation protocol (Harris et al. 1978 Biochem. 17:3251-3256; Galau et al. 1981 J. Biol. Chem. 256:2551-2560). Poly A+ RNA was isolated using oligo dT column chromatography according to manufacturers' protocols (PHARMACIA or

STRATAGENE) or using oligotex-dT latex particles (QIAGEN).

Construction of tissue-specific cDNA libraries

Flower, one day silique, three day silique, leaf, root, and developing seed cDNA libraries were each constructed from 5 µg poly A+ RN using the ZAP cDNA synthesis kit (Stratagene). cDNAs were directionally cloned into the EcoRI and XhoI sites of pBluescript SK(-) in the λ -ZAPII vector (Short et al. 1988 Nucleic Acids Res. 16:7583-7600). Nonrecombinant phage plaques were identified by blue color development on NZY plates containing X-gal (5 bromo-4chloro-3-indoyl- β -D-galactopyranoside) and IPTG (isopropyl-1-thio- β -D-galactopyranoside). The nonrecombinant backgrounds for the flower, one day silique, three day silique, leaf, root, and developing seed cDNA libraries were 2.8%, 2%m 3.3%, 6.5%, 2.5%, and 1.9% respectively.

Random priming DNA labeling

The cDNA inserts of isolated clones (unhybridized cDNAs) were excised by EcoRI/XhoI double digestion and gel-purified for random priming labeling. Klenow reaction mixture contained 50 ng DNA templates, 10 mM Tris-HCl, pH 7.5, 5 mM MgCl₂, 7.5 mM DTT, 50 uM each of dCTP, dGTP, and dTTP, 10 uM hexamer random primbers (Boehringer Mannheim), 50 μ Ci α -32 P-dATP, 3000 Ci/mmole, 10 mCi/ml (DuPont), and 5 units of DNA polymerase I Klenow fragment (New England

Biolabs). The reactions were carried out at 37°C for one hour. Aliquots of diluted reaction mixtures were used for TCA precipitation and alkaline denaturing gel analysis. Hybridization probes were labeled only with Klenow DNA polymerase and the unincorporated dNTPs were removed using Sephadex R G-50 spin columns (Boehringer Mannheim).

Random Primed PCR

Double-stranded cDNA was synthesized from poly A+ RNA isolated from Arabidopsis root tissue using the cDNA Synthesis System (GIBCO BRL) with oligo dT12-18 as primers. cDNAs longer than 300 bp were enriched by Sephacryl S-400 column chromatography (Stratagene). Fractionated cDNAs were used as templates for RP-PCR labeling. The reaction contained 10 mM Tris-HCl, ph 9.0, 50 mM KCl, 0.1% Triton X-100, 2 mM·MgCl2, 5 units Taq DNA polymeras (PROMEGA), 200 μM dCTP, cGTP, and dTTP, and different concentrations of hexamer random primers α -32P dATP, 800 mCi/mmole, 10 mCi/ml (DuPont), and cold dATP in a final volume of 25 μl. After an initial 5 minutes at 95°C, different reactions were run through different programs to optimize RP-PCR cDNA conditions. Unless otherwise indicated, the following program was used for most RP-PCR cDNA probe labeling: 95°C/5 minutes, then 40 cycles of 95°C 30 seconds, 18°C/1 second, ramp to 30°C at a rate of 0.1°C/second. 72°C/1 minute. products were phenol/chloroform extracted and ethanol

precipitated or purified by passing through Sephadex G-50 spin columns (Boehringer Mannheim).

Clone blot virtual subtraction

Mass excision of λ-ZAP cDNA libraries was carried out by co-infecting XL1-Blue MRF' host cells with recombinant phage from the libraries and ExAssist helper phage (STRATAGENE). Excised phagemids were rescued by SOLR cells. Plasmid DNAs were prepared by boiling mini-prep method (Holmes et al. 1981 Anal. Biochem. 114:193-197) from randomly isolated clones. cDNA inserts were excised by EcoRI and XhoI double digestion, and resolved on 1% agarose gels. were denatured in 0.5 N NaOH and 1.5 m NaCl for 45 minutes, neutralized in 0.5 M Tris-HCl, pH 8.0, and 1.5 M NaCl for 45 minutes, and then transferred by blotting to nylon membranes (Micron Separations, Inc.) in 10X SSC overnight. After one hour prehybridization at 65°C, root RP-cDNA probe was added to the same hybridization buffer containing 1% bovine albumin fraction V (Sigma), 1 mM EDTA, 0.5 M NaHPO4, pH 7.2, The hybridization continued for 24 hours at 7% SDS. 65°C. The filters were washed in 0.5% bovine albumin, 1 mM EDTA, 40 mM NaHPO4, pH 7.2, 5% SDS for ten minutes at room temperature, and 3 x 10 minutes in 1 mM EDTA, 40 mM NaHPO4, pH 7.2, 1% SDS at 65°C. Autoradiographs were exposed to X-ray films (Kodak) for two to five days at -80°C.

Hybridization of resulting blots with root RP-PCR probes "virtually subtracted" seed cDNAs shared

with the root mRNA population. The remaining seed cDNAs representing putative seed-specific cDNAs, including those encoding oleosins, were sequenced by the cycle sequencing method, thereby identifying AtS21 as an oleosin cDNA clone.

Sequence analysis of AtS21

The oleosin cDNA is 834 bp long including an 18 bp long poly A tail (Fig. 4, SEQ ID NO:2) high homology to other oleosin genes from Arabidopsis as well as from other species. Recently, an identical oleosin gene has been reported (Zou, et al., 1996, Plant Mol. Biol. 31:429-433). The predicted protein is 191 amino acids long with a highly hydrophobic middle domain flanked by a hydrophilic domain on each side. The existence of two upstream in frame stop codons and the similarity to other oleosin genes indicate that this cDNA is full-length. Since there are two in frame stop codons just upstream of the first ATG, this cDNA is considered to be a full length cDNA (Figure 4, SEQ ID NO:2). The predicted protein has three distinctive domains based on the distribution of its amino acid residues. Both the N-terminal and C-terminal domains are rich in charged residues while the central domain is absolutely hydrophobic (Figure 5). As many as 20 leucine residues are located in the central domain and arranged as repeats with one leucine occurring every 7-10 residues. Other non-polar amino acid residues are also clustered in the central domain making this domain absolutely hydrophobic (Figure 6).

Extensive searches of different databases using both AtS21 cDNA and its predicted protein sequence identified oleosins from carrot, maize, cotton, rapeseed, Arabidopsis, and other plant species. The homology is mainly restricted to the central hydrophobic domain. Seven Arabidopsis oleosin sequences were found. AtS21 represents the same gene as Z54164 which has a few more bases in the 5' untranslated region. The seven Arabidopsis oleosin sequences available so far were aligned to each other (Figure 7). The result suggested that the seven sequences fall into three groups. The first group includes AtS21 (SEQ ID NO:5), X91918 (SEQ ID NO:6), and the partial sequence Z29859 (SEQ ID NO:7). Since X91918 (SEQ ID NO:6) has only its last residue different from AtS21 (SEQ ID NO:5), and since Z29859 (SEQ ID NO:7) has only three amino acid residues which are different from AtS21 (SEQ ID NO:5), all three sequences likely represent the same gene. The two sequences of the second group, X62352 (SEQ ID NO:8) and Atol3 (SEQ ID NO:9), are different in both sequence and length. Thus, there is no doubt that they represent two independent genes. Like the first group, the two sequences of the third group, X91956 (SEO ID NO:10) and L40954 (SEO ID NO:11), also have only three divergent residues which may be due to sequence errors. Thus, X91956 (SEQ ID NO:10) and L40954 (SEQ ID NO:11) likely represent the same gene. Unlike all the other oleosin sequences which were predicted from cDNA sequences, X62352 (SEQ ID NO:8)

was deduced from a genomic sequence (Van Rooigen et al. 1992 Plant Mol. Biol. 18:1177-1179). In conclusion, four different Arabidopsis oleosin genes have been identified so far, and they are conserved only in the middle of the hydrophobic domain.

Northern Analysis

In order to characterize the expression pattern of the native AtS21 gene, Northern analysis was performed as described in Example 4 except that the probe was the AtS21 cDNA (pAN1 insert) labeled with 12 P-dATP to a specific activity of 5 x 10^{8} cpm/ug.

Results indicated that the AtS21 gene is strongly expressed in developing seeds and weakly expressed in silique coats (Figure 8A). A much larger transcript, which might represent unprocessed AtS21 pre-mRNA, was also detected in developing seed RNA. AtS21 was not detected in flower; leaf, root (Figure 8A), or one day silique RNAs. A different Northern analysis revealed that AtS21 is also strongly expressed in imbibed germinating seeds (Figs. 13A and 13B)

EXAMPLE 6

Characterization of Oleosin
Genomic Clones and Isolation of Oleosin Promoter

Genomic clones were isolated by screening an Arabidopsis genomic DNA library using the full length cDNA (AtS21) as a probe. Two genomic clones were mapped by restriction enzyme digestion followed by Southern hybridization using the 5' half of the cDNA cleaved by SacI as a probe. A 2 kb SacI fragment was subcloned and sequenced (Fig. 9, SEQ ID NO:35). Two regions of the genomic clone are identical to the cDNA sequence. A 395 bp intron separates the two regions.

The copy number of AtS21 gene in the Arabidopsis genome was determined by genomic DNA Southern hybridization following digestion with the enzymes BamHI, EcoRI, HindIII, SacI and XbaI, using the full length cDNA as a probe (Figure 8B). A single band was detected in all the lanes except SacI digestion where two bands were detected. Since the cDNA probe has an internal SacI site, these results indicated that AtS21 is a single copy gene in the Arabidopsis genome. Since it has been known that Arabidopsis genome contains different isoforms of oleosin genes, this Southern analysis also demonstrates that the different oleosin isoforms of Arabidopsis are divergent at the DNA sequence level.

Two regions, separated by a 395 bp intron, of the genomic DNA fragment are identical to AtS21 cDNA sequence. Database searches using the 5' promoter sequence upstream of AtS21 cDNA sequence did

not identify any sequence with significant homology. Furthermore, the comparison of AtS21 promoter sequence with another *Arabidopsis* oleosin promoter isolated previously (Van Rooijen, et al., 1992) revealed little similarity. The AtS21 promoter sequence is rich in A/T bases, and contains as many as 44 direct repeats ranging from 10 bp to 14 bp with only one mismatch allowed. Two 14 bp direct repeats, and a putative ABA response element are underlined in Figure 9.

EXAMPLE 7

Construction of AtS21
Promoter/GUS Gene Expression Cassette and Expression
Patterns in Transgenic Arabidopsis and Tobacco

Construction of AtS21 promoter/GUS gene expression cassette

The 1267 bp promoter fragment starting from the first G upstream of the ATG codon of the genomic DNA fragment was amplified using PCR and fused to the GUS reporter gene for analysis of its activity. The promoter fragment of the AtS21 genomic clone was amplified by PCR using the T7 primer GTAATACGACTCACTATAGGGC (SEQ ID NO:13) and the 21P primer GGGGATCCTATACTAAAACTATAGAGTAAAGG (SEQ ID NO:14) complementary to the 5' untranslated region upstream of the first ATG codon (Figure 9). A BamHI cloning site was introduced by the 21P primer. The amplified fragment was cloned into the BamHI and SacI sites of pBluescript KS (Stratagene). Individual clones were sequenced to check possible PCR mutations as well as the orientation of their inserts. The correct clone was digested with BamHI and HindIII, and the excised promoter fragment (1.3 kb) was cloned into the corresponding sites of pBI101.1 (Jefferson, R.A. 1987a, Plant Mol. Biol. Rep. 5:387-405; Jefferson et al., 1987b, EMBO J. 6:3901-3907) upstream of the GUS The resultant plasmid was designated pAN5 (Fig. The AtS21 promoter/GUS construct (pAN5) was introduced into both tobacco (by the leaf disc method, Horsch et al., 1985; Bogue et al. 1990 Mol. Gen. Gen.

221:49-57) and Arabidopsis Colombia ecotype via vacuum infiltration as described by Bechtold, et al. C.R. Acad. Sci. Paris, 316:1194-1199. Seeds were sterilized and selected on media containing 50 μ g/ml kanamycin, 500 μ g/ml carbenicillin. GUS activity assay: Expression patterns of the reporter GUS gene were revealed by histochemical staining (Jefferson, et al., 1987a, Plant Mol. Biol. Rep. 5:387-405). Different tissues were stained in substrate solution containing 2 mg/ml 5-bromo-4chloro-3-indolyl-β-D-glucuronic acid (X-Gluc) (Research Organics, Inc.), 0.5 mM potassium ferrocyanide, and 0.5 mM potassium ferricyanide in 50 mM sodium phosphate buffer, pH 7.0 at 37°C overnight, and then dehydrated successively in 20%, 40% and 80% ethanol (Jefferson, et al., 1987). Photographs were taken using an Axiophot (Zeiss) compound microscope or Olympus SZH10 dissecting microscope. Slides were converted to digital images using a Spring/Scan 35LE slide scanner (Polaroid) and compiled using Adobe Photoshop™ 3.0.5 and Canvas™ 3.5.

GUS activities were quantitatively measured by fluorometry using 2 mM 4-MUG (4-methylumbelliferyl- $\beta\text{-}D\text{-}glucuronide)$ as substrate (Jefferson, et al., 1987). Developing Arabidopsis seeds were staged according to their colors, and other plant tissues were collected and kept at -80°C until use. Plant tissues were ground in extraction buffer containing 50 mM sodium phosphate, pH 7.0, 10 mM EDTA, 10 mM $\beta\text{-}$ mercaptoethanol, 0.1% Triton X-100, and 0.1% sodium

lauryl sarcosine. The tissue debris was removed by 5 minutes centrifugation in a microfuge. The supernatant was aliquoted and mixed with substrate and incubated at 37°C for 1 hour. Three replicas were assayed for each sample. The reactions were stopped by adding 4 volumes of 0.2 M sodium carbonate. Fluorescence was read using a TKO-100 DNA fluorometer (Hoefer Scientific Instruments). Protein concentrations of the extracts were determined by the Bradford method (Bio Rad).

Expression patterns of AtS21 promoter/GUS in transgenic Arabidopsis and tobacco

In Arabidopsis, GUS activity was detected in green seeds, and node regions where siliques, cauline leaves and branches join the inflorescence stem (Figures 11A and 11B). No GUS activity was detected in any leaf, root, flower, silique coat, or the internode regions of the inflorescence stem. Detailed studies of the GUS expression in developing seeds revealed that the AtS21 promoter was only active in green seeds in which the embryos had already developed beyond heart stage (Figures 11C and 11G). youngest embryos showing GUS activity that could be detected by histochemical staining were at early torpedo stage. Interestingly, the staining was only restricted to the lower part of the embryo including hypocotyl and embryonic radical. No staining was detected in the young cotyledons (Figures 11D and 11E). Cotyledons began to be stained when the embryos

were at late torpedo or even early cotyledon stage (Figure 11F and 11H). Later, the entire embryos were stained, and the staining became more intense as the embryos matured (Figures 11I and 11J). It was also observed that GUS gene expression was restricted to the embryos. Seed coat and young endosperm were not stained (Figure 11C).

GUS activity was also detected in developing seedlings. Young seedlings of 3-5 days old were stained everywhere. Although some root hairs close to the hypocotyl were stained (Figure 11K), most of the newly formed structures such as root hairs, lateral root primordia and shoot apex were not stained (Figures 11L and 11N). Later, the staining was restricted to cotyledons and hypocotyls when lateral roots grew from the elongating embryonic root. The staining on embryonic roots disappeared. No staining was observed on newly formed lateral roots, true leaves nor trichomes on true leaves (Figures 11M and 11N).

AtS21 promoter/GUS expression patterns in tobacco are basically the same as in Arabidopsis. GUS activity was only detected in late stage seeds and different node regions of mature plants. In germinating seeds, strong staining was detected throughout the entire embryos as soon as one hour after they were dissected from imbibed seeds. Mature endosperm, which Arabidopsis seeds do not have, but not seed coat was also stained (Figure 12A). The root tips of some young seedlings of one transgenic line

were not stained (Figure 12B). Otherwise, GUS expression patterns in developing tobacco seedlings were the same as in Arabidopsis seedlings (Figures 12B, 12C, and 12D). Newly formed structures such as lateral roots and true leaves were not stained.

AtS21 mRNA levels in developing seedlings

Since the observed strong activities of AtS21 promoter/GUS in both Arabidopsis and tobacco seedlings are not consistent with the seed-specific expression of oleosin genes, Northern analysis was carried out to determine if AtS21 mRNA was present in developing seedlings where the GUS activity was so strong. RNAs prepared from seedlings at different stages from 24 hours to 12 days were analyzed by Northern hybridization using AtS21 cDNA as the probe. Surprisingly, AtS21 mRNA was detected at a high level comparable to that in developing seeds in 24-48 hour The mRNA level dropped dramatically imbibed seeds. when young seedlings first emerged at 74 hours (Figures 13A and 13B). In 96 hour and older seedlings, no signal was detected even with a longer exposure (Figure 13B). The loadings of RNA samples were checked by hybridizing the same blot with a tubulin gene probe (Figure 13C) which was isolated and identified by EST analysis as described in Example 2. Since AtS21 mRNA was so abundant in seeds, residual AtS21 probes remained on the blot even after extensive These results indicated that AtS21 mRNA detected in imbibed seeds and very young seedlings are the carry-over of AtS21 mRNA from dry seeds. It has recently been reported that an oleosin Ato12 mRNA (identical to AtS21) is most abundant in dry seeds (Kirik, et al., 1996 Plant Mol. Biol. 31(2):413-417.) Similarly, the strong GUS activities in seedlings were most likely due to the carry-over of both β -glucuronidase protein and the de novo synthesis of β -glucuronidase from its mRNA carried over from the dry seed stage.

EXAMPLE 8

Activity comparison between the AtS21 promoter and the 35S promoter

The GUS activities in transgenic Arabidopsis developing seeds expressed by the AtS21 promoter were compared with those expressed by the 35S promoter in the construct pBI221 (Jefferson et al. EMBO J. 6:3901-The seeds were staged according to their colors (Table 2). The earliest stage was from globular to late heart stage when the seeds were still white but large enough to be dissected from the siliques. AtS21 promoter activity was detected at a level about three times lower than that of the 35S promoter at this stage. 35S promoter activity remained at the same low level throughout the entire embryo development. In contrast, AtS21 promoter activity increased quickly as the embryos passed torpedo stage and reached the highest level of 25.25 pmole 4-MU/min. μ g protein at mature stage (Figure 5-The peak activity of the AtS21 promoter is as much as 210 times higher than its lowest activity at globular to heart stage, and is close to 100 times higher than the 35S promoter activity at the same stage (Table 2). The activity levels of the AtS21 promoter are similar to those of another Arabidopsis oleosin promoter expressed in Brassica napus (Plant et al. 1994, Plant mol. Biol. 25:193-205. AtS21 promoter activity was also detected at background level in leaf. The high standard deviation, higher than the average itself, indicated that the GUS activity was

only detected in the leaves of some lines (Table 2). On the other hand, 35S promoter activity in leaf was more than 20 times higher than that in seed. The side by side comparisons of activities between AtS21 promoter and 35S promoter is shown in Figure 14.

Although the AtS21 promoter activity was about 3 times lower in dry seed of tobacco than in Arabidopsis dry seed, the absolute GUS activity was still higher than that expressed by the 35S promoter in Arabidopsis leaf (Table 2). No detectable AtS21 promoter activity was observed in tobacco leaf (Figure 14).

Comparison of the AtS21 promoter versus the 35S promoter revealed that the latter is not a good promoter to express genes at high levels in developing seeds. Because of its consistent low activities throughout the entire embryo development period, 35S promoter is useful for consistent low level expression of target genes. On the other hand, the AtS21 promoter is a very strong promoter that can be used to express genes starting from heart stage embryos and accumulating until the dry seed stage. The 35S promoter, although not efficient, is better than the AtS21 promoter in expressing genes in embryos prior to heart stage.

			LEA	0.08±0. 6.56±0. 0.01±0.
1			BROWN DRY SEED	24.38±10.85 0.31±0.02 8.81±0.21
5		STRUCTS	green/yellow/brown mature	25.25±4.64 0.26±0.04
10 .		OTER/GUS CON	DARK GREEN GR LATE C	21.85±4.45 0.33±0.06
15	TABLE 2	OF AtS21 and 35S PROMOTER/GUS CONSTRUCTS	LIGHT GREEN EARLY C	18.99±3.75 0.28±0.03
20	•	OF AtS21	YELLOW T-C	6.77±1.25 0.29±0.04
25		GUS ACTIVITIES	WHITE/YELLOW H-T	0.12±0.17 1.35±1.57 0.30±0.06 0.25±0.08 (In tobacco)
30		าย	WHITE G·H	0.12±0.17 0.30±0.06 (In tobacco)
35			COLOR STAGE	AtS21 358 AtS21

Abbreviations: G, globular stage; H, heart stage; T, torpedo stage; C, cotyledon stage. The GUS activities are in pmole 4-MU/µg protein.min. For AtS21 promoter the numbers are the average of five independent lines with standard deviations. Three repeats were assayed for each line. For 35S promoter the numbers are the average of three repeats of the same line with standard deviations.

EXAMPLE 9

Expression of the Borage Δ^6 -Desaturase Gene Under the Control of the AtS21 Promoter and Comparison to Expression Under the Control of the CaMV 35S Promoter

In order to create an expression construct with the AtS21 promoter driving expression of the borage $\Delta 6$ -desaturase gene, the GUS coding fragment from pAN5 was removed by digestion with SmaI and ECOICR I. The cDNA insert of pAN1 (Example 2) was then excised by first digesting with XhoI (and filling in the residual overhang as above), and then digesting with SmaI. The resulting fragment was used to replace the excised portion of pAN5, yielding pAN3.

After transformation of tobacco and Arabidopsis following the methods of Example 7, levels of Δ^6 -desaturase activity were monitored by assaying the corresponding fatty acid methyl esters of its reaction products, γ -linolenic acid (GLA) and octadecatetraenoic acid (OTA) using the methods referred to in Example 3. The GLA and OTA levels (Table 3) of the transgenic seeds ranged up to 6.7% of C18 fatty acids (Mean = 3.1%) and 2.8% (Mean = 1.1%), respectively. No GLA or OTA was detected in the leaves of these plants. In comparison, CaMV 35 S promoter/ Δ^6 -desaturase transgenic plants produced GLA levels in seeds ranging up to 3.1% of C18 fatty acids (Mean = 1.3%) and no measurable OTA in seeds.

EXPRESSION OF THE BORAGE A *- DESATURASE IN TRANSGENIC PLANTS

PROMOTER	PLANT		SEED	А			ជ	LEAP	
		GLA*	RANGE	OTA*	RANGE	et.	RANGE	OTA	ota range
Cauliflower mosaic virus 35S	tobacco	1.3	0.7-3.1	n. d		50	19-22	9.7 8-11	8-11
Arabidopsis oleosin	Arabidopsis 3.1	3.1	0-6.7	1.1	1.1 0-2.8 n.d.	n.d.		n.d	

*mean value expressed as the percent of the C_{1s} fatty acids n.d. not detected

EXAMPLE 10

Transformation of Oilseed Rape With an Expression Cassette Which Comprises the Oleosin 5' Regulatory Region Linked to the Borage Delta 6-Desaturase Gene

Oilseed rape, Cv. Westar, was transformed with the strain of Agrobacterium tumefaciens EHA105 containing the plasmid pAN3 (i.e. the borage $\Delta 6$ -desaturase gene under the control of the Arabidopsis oleosin promoter-Example 9).

Terminal internodes of Westar were cocultivated for 2-3 days with induced Agrobacterium
tumefaciens strain EHA105 (Alt-Moerbe et al. 1988 Mol.
Gen. Genet. 213:1-8; James et al. 1993 Plant Cell
Reports 12:559-563), then transferred onto
regeneration medium (Boulter etal. 1990 Plant Science
70:91-99; Fry et al. 1987 Plant Cell Reports 6:321325). The regenerated shoots were transferred to
growth medium (Pelletier et al. 1983 Mol.Gen. Menet.
191:244-250), and a polymerase chain reaction (PCR)
test was performed on leaf fragments to assess the
presence of the gene.

DNA was isolated from the leaves according to the protocol of KM Haymes et al. (1996) Plant Molecular Biology Reporter 14(3):280-284, and resuspended in 100µl of water, without RNase treatment. 5µl of extract were used for the PCR reaction, in a final volume of 50µl. The reaction was performed in a Perkin-Elmer 9600 thermocycler, with the following cycles:

1 cycle: 95°C, 5 minutes

30 cycles: 95°C, 45 sec; 52°C, 45 sec

72°C, 1 minute

1 cycle: 72°C, 5 minutes

and the following primers (derived from near the metal box regions, as indicated in Fig. 1, SEQ. NO.:1):

5' TGG AAA TGG AAC CAT AA 3'

5' GGA AAC AAA TGA TGC TC 3'

Amplification of the DNA revealed the expected 549 base pair PCR fragment (Figure 17).

The positive shoots were transferred to elongation medium, then to rooting medium (DeBlock et al 1989 Plant Physiol. 91:694-701). Shoots with a well-developed root system were transferred to the greenhouse. When plants were well developed, leaves were collected for Southern analysis in order to assess gene copy number.

Genomic DNA was extracted according to the procedure of Bouchez et al. (1996) Plant Molecular Biology Reporter 14:115-123, digested with the restriction enzymes Bgl I and/or Cla I, electrophoretically separated on agarose gel (Maniatis et al. 1982, in Molecular Cloning; a Laboratory Manual. Cold Spring Harbor Laboratory Press, Cold Spring Harbor/NY), and prepared for transfer to nylon membranes (Nytran membrane, Schleicher & Schuell) according to the instructions of the manufacturer. DNA was then transferred to membranes overnight by capillary action using 20XSSC (Maniatis et al. 1982).

Following transfer, the membranes were crosslinked by UV (Stratagene) for 30 seconds and pre-hybridized for 1 hour at 65°C in 15 ml of a solution containing 6XSSC, 0.5%SDS and 2.25% w/w dehydrated skim milk in glass vials in hybridization oven (Appligene). The membranes were hybridized overnight in the same solution containing a denatured hybridization probe radiolabelled with 32P to a specific activity of 108 cpm/µg by the random primer method (with the Ready-To-Go kit obtained from Pharmacia). The probe represents a PCR fragment of the borage delta 6-desaturase gene (obtained in the conditions and with the primers detailed above). After hybridization, the filters were washed at 65°C in 2XSSC, 0.1% SDS for 15 minutes, and 0.2XSSC, 0.1%SDS for 15 minutes. The membranes were then wrapped in Saran-Wrap and exposed to Kodak XAR film using an intensifying screen at -70°C in a light-proof cassette. Exposure time was generally 3 days.

The results obtained confirm the presence of the gene. According to the gene construct, the number of bands in each lane of DNA digested by *Bgl* I or *Cla* I represents the number of delta 6-desaturase genes present in the genomic DNA of the plant. The digestion with *Bgl* 1 and *Cla* 1 together generates a fragment of 3435 bp.

The term "comprises" or "comprising" is defined as specifying the presence of the stated features, integers, steps, or components as referred to in the claims, but does not preclude the presence or addition of one or more other features, integers, steps, components, or groups thereof.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Rhone Poulenc Agro
 Thomas, Terry L.
 Li, Zhongsen
- (ii) TITLE OF INVENTION: AN OLEOSIN 5' REGULATORY REGION FOR THE MODIFICATION OF PLANT SEED LIPID COMPOSITION
- (iii) NUMBER OF SEQUENCES: 35
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Scully, Scott, Murphy & Presser
 - (B) STREET: 400 Garden City Plaza
 - (C) CITY: Garden City
 - (D) STATE: New York
 - (E) COUNTRY: USA
 - (F) ZIP: 11530
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: 08/831,575
 - (B) FILING DATE: 9 April 1997
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: DiGiglio, Frank S.
 - (B) REGISTRATION NUMBER: 31,346
 - (C) REFERENCE/DOCKET NUMBER: 10203
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (516) 742-4343
 - (B) TELEFAX: (516) 742-4366
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1684 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 43..1387

-59-

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

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5	2,3	bys	Tyr	116	10	ser	Asp	GIU	Leu	Lys 15	Asn	His	Asp	Lys	CCC Pro 20		102
Cly	nop	Deu	пр	25	ser	116	GIN	GIY	30	Ala	Tyr	Asp	Val	Ser 35			150
	val	шуз	40	ms	·	GIA	GIÀ	ser 45	Pne	Pro	Leu	Lys	Ser 50	Leu	GCT Ala		198
0	0111	55	Vai	1111	wsp	VIG	60	vaı	ATa	Phe	His	Pro 65	Ala	Ser			246
	70	ASII	beu	nsp	гÀ2	75	Pne	Thr	GIA	TAT Tyr	Tyr 80	Leu	Lys	Asp	Tyr		294
85	V	·	Gru	vaı	90	ьуs	Asp	Tyr	Arg	AAG Lys 95	Leu	Val	Phe	Glu	Phe 100		342
001	БуЗ	Mec	GLY	105	Tyl	ASp	rys	гÀЗ	110	CAT His	Ile	Met	Phe	Ala 115	Thr		390
TTG Leu	TGC Cys	TTT Phe	ATA Ile 120	GCA Ala	ATG Met	CTG Leu	TTT Phe	GCT Ala 125	λTG Met	AGT Ser	GTT Val	TAT Tyr	GGG Gly 130	GTT Val	TTG Leu		438
TTT Phe	TGT Cys	GAG Glu 135	GGT Gly	GTT Val	TTG Leu	GTA Val	CAT His 140	TTG Leu	TTT Phe	TCT Ser	GGG Gly	TGT Cys 145	TTG Leu	ATG Met	GGG Gly		486
TTT Phe	CTT Leu 150	TGG Trp	ATT Ile	CAG Gln	AGT Ser	GGT Gly 155	TGG Trp	ATT Ile	GGA Gly	CAT His	GAT Asp 160	GCT Ala	GGG Gly	CAT His	TAT Tyr		534
ATG Met 165	GTA Val	GTG Val	TCT Ser	GAT Asp	TCA Ser 170	AGG Arg	CTT Leu	AAT Asn	AAG Lys	TTT Phe 175	ATG Met	GGT Gly	ATT Ile	TTT Phe	GCT Ala 180		582
GCA Ala	AAT Asn	TGT Cys	CTT Leu	TCA Ser 185	GGA Gly	ATA Ile	AGT Ser	пе	GGT Gly 190	TGG Trp	TGG Trp	AAA Lys	TGG Trp	AAC Asn 195	CAT His	•	630

AAT Asn	GCA Ala	CAT His	CAC His 200	ATT Ile	GCC Ala	TGT Cys	AAT Asn	AGC Ser 205	CTT Leu	GAA Glu	TAT Tyr	GAC Asp	CCT Pro 210	GAT Asp	TTA	678
GIN	туг	215	Pro	Pne	Leu	vai	Va1 220	Ser	Ser	Lys	Phe	Phe 225	Gly	Ser	CTC Leu	726
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245	File	Val	ser	туг	CAA Gln 250	HIS	Trp	Thr	Phe	Tyr 255	Pro	Ile	Met	Cys	Ala 260	822
ALG	nrg	Deu	ASII	265	TAT Tyr	val	GIN	ser	270	Ile	Met	Leu	Leu	Thr 275	Lys	870
ALG	ASII	vai	280	TŸĽ	CGA Arg	Ala	GIn	G1u 285	Leu	Leu	Gly	Cys	Leu 290	Val	Phe	918
Ser	116	295	ıyr	Pro	TTG Leu	Leu	300	Ser	Cys	Leu	Pro	Asn 305	Trp	Gly	Glu	966
nrg	310	Mer	rne	vaı		315	Ser	Leu	Ser	Val	Thr 320	Gly	Met	Gln	Gln ·	1014
325	GIII	rne	ser	Leu	AAC Asn 330	HIS	Pne	Ser	Ser	335	Val	Tyr	Val	Gly	Lys 340	1062
	nys.	GIÀ	·ASII	345	TGG Trp	Pne	Giu	Lys	GIn 350	Thr	Asp	Gly	Thr	Leu 355	Asp	1110
116	ser	Cys	360	Pro	TGG Trp	мес	Asp	Trp 365	Phe	His	Gly	Gly	Leu 370	Gln	Phe	1158
GIII	rre	375	HIS	HIS	TTG Leu	Phe	Pro 380	Lys	Met	Pro	Arg	Cys 385	Asn	Leu	Arg	1206
ьуs	390	ser	Pro	Tyr	GTG Val	395	Glu	Leu	Cys	Lys	Lys 400	His	Asn	Leu	Pro	1254
TAC Tyr 405	AAT Asn	TAT Tyr	GCA Ala	TCT Ser	TTC Phe 410	TCC Ser	AAG Lys	GCC Ala	Asn	GAA Glu 415	atg Met	ACA Thr	CTC Leu	Arg	ACA Thr 420	1302

-614

TT	ı yığ	G AA(g Asi	C ACA	A GC2 c Ala 425	T TIE	G CAC	G GCT n Ala	r AG	G GA' g Ası 430	5 TT	A ACC	C AAC	G CC	G CTO	C CCG u Pro	;	1350
λA(Lys	G AAT G AST	r TT(1 Lei	GT7 1 Val 440		GAN Glu	A GCT 1 Ala	CT Lei	CAC 1 His 445	s Thi	r CAT	r GG	r T /	እ እአለ	ITAC	CC		1397
															rggtt		1457
															TATAT		1517
															\ATTG'		1577
GTO	CTCA	λΤλ	TCTG	TATA	TT 1	'GGAA	TGT	C TI	TTGT	ССУС	GTC	GTT	TCA	GTTC	BANGC'	TC	1637
ATC	TGTA	CTT	CTAT	'AGAC	TT 1	GTTI	'AAA'	'G G'I	TATO	TCAT	GTI	'ATTI	ŗ				1684
(2)	INF	ORMA	TION	FOR	SEO	TD	พดงว										
			SEQU (A (B	ENCE) LE) TY	CHA NGTH PE:		ERIS 5 am o ac	TICS ino id	: acid	s							
	(ii)	MOLE														
			SEQU:						O ID	NO.	2.	•					
	λla	λla	Gln	Ile 5	Lys	Lys	Туr	Ile	Thr 10	Ser	Asp		Leu	15			
			- •					23					Lys 30				
Asp	Val	Ser 35	Asp	Trp	Val	Lys	Asp	His	Pro	Gly	Gly	Şer	Phe	Pro	Leu		
Lys	Ser 50	Leu	Ala	Gly	Gln	Glu 55	Val	Thr	Asp	Ala	Phe 60	45 Val	Ala	Phe	His		
Pro 65	λla	Ser	Thr	Trp	Lys 70	λsn	Leu	Asp	Lys	Phe 75	Phe	Thr	Gly	Tyr	Tyr 80		
Leu	Lys	Asp	Tyr	Ser 85	Val	Ser	Glu	Val	Ser 90	Lys	Asp	Tyr	Arg	Lys 95	Leu		
Val	Phe	Glu	Phe 100	Ser	Lys	Met	Gly	Leu 105	Tyr	Asp	Lys	Lys	Gly 110	His	Ile		
Met	Phe	λla 115	Thr	Leu	Суѕ	Phe	Ile 120	Ala	Met	Leu	Phe	Ala 125	Met	Ser	Val		
Tyr	Gly 130	Val	Leu	Phe	Cys	Glu 135							•				

(2) INFORMATION	FOR	SEQ	ID	NO:	3	:
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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 834 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: double
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: CDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 31..603
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

								_									
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CGT Arg	GTG Val 145	CAT His	GTA Val	GAC Asp	CGT Arg	ACT Thr 150	GAC Asp	AAA Lys	CGT Arg	GTT Val	CAT His 155	CAG Gln	CCA Pro	AAC Asn	TAC Tyr	1	02
GAA Glu 160	GAT Asp	GAT Asp	GTC Val	GGT Gly	TTT Phe 165	GGT Gly	GGC Gly	ТАТ Туг	GGC Gly	GGT G1y 170	TAT Tyr	GGT Gly	GCT Ala	GGT Gly	TCT Ser 175	1!	50
GAT Asp	TAT Tyr	AAG Lys	AGT Ser	CGC Arg 180	GGC Gly	CCC Pro	TCC Ser	ACT Thr	AAC Asn 185	CAA Gln	ATC Ile	TTG Leu	GCA Ala	CTT Leu 190	ATA Ile	19	98
GCA Ala	GGA Gly	GTT Val	CCC Pro 195	ATT Ile	GGT Gly	GGC Gly	ACA Thr	CTG Leu 200	CTA Leu	ACC Thr	CTA Leu	GCT Ala	GGA Gly 205	CTC Leu	ACT Thr	24	46
CTA Leu	GCC Ala	GGT Gly 210	TCG Ser	GTG Val	ATC Ile	GGC Gly	TTG Leu 215	CTA Leu	GTC Val	TCC Ser	ATA Ile	CCC Pro 220	CTC Leu	TTC Phe	CTC Leu	29	94
CTC Leu	TTC Phe 225	AGT Ser	CCG Pro	GTG Val	ATA Ile	GTC Val 230	CCG Pro	GCG Ala	GCT Ala	CTC Leu	ACT Thr 235	ATT Ile	GGG Gly	CTT Leu	GCT Ala	34	42
GTG Val 240	ACG Thr	GGA Gly	ATC Ile	TTG Leu	GCT Ala 245	TCT Ser	GGT Gly	TTG Leu	TTT Phe	GGG Gly 250	TTG Leu	ACG Thr	GGT Gly	CTG Leu	AGC Ser 255	39	90
TCG Ser	GTC Val	TCG Ser	TGG Trp	GTC Val 260	CTC Leu	AAC Asn	TAC Tyr	CTC Leu	CGT Arg 265	Gly	ACG Thr	AGT Ser	GAT Asp	ACA Thr 270	GTG Val	43	38

CCA GAG CAA TTG GAC TAC GCT AAA CGG CGT ATG GCT GAT GCG GTA GGC Pro Glu Gln Leu Asp Tyr Ala Lys Arg Arg Met Ala Asp Ala Val Gly 275 280 285	486
TAT GCT GGT ATG AAG GGA AAA GAG ATG GGT CAG TAT GTG CAA GAT AAG Tyr Ala Gly Met Lys Gly Lys Glu Met Gly Gln Tyr Val Gln Asp Lys 290 295 300	534
GCT CAT GAG GCT CGT GAG ACT GAG TTC ATG ACT GAG ACC CAT GAG CCG Ala His Glu Ala Arg Glu Thr Glu Phe Met Thr Glu Thr His Glu Pro 305 310 315	. 582
GGT AAG GCC AGG AGA GGC TCA TAAGCTAATA TAAATTGCGG GAGTCAGTTG Gly Lys Ala Arg Arg Gly Ser 320 325	633
GAAACGCGAT AAATGTAGTT TTACTTTTAT GTCCCAGTTT CTTTCCTCTT TTAAGAATAT	693
CTTTGTCTAT ATATGTGTTC GTTCGTTTTG TCTTGTCCAA ATAAAAATCC TTGTTAGTGA	753
ANTANGAAAT GAAATAAATA TGTTTTCTTT TTTGAGATAA CCAGAAATCT CATACTATTT	813
ТСТАЛАЛЛАА ААЛААЛАЛА А	
(2) INFORMATION FOR SEQ ID NO:4:	834
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 191 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: protein	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
Met Ala Asn Val Asp Arg Asp Arg Val His Val Asp Arg Thr Asp 1 5 10 15	
Lys Arg Val His Gln Pro Asn Tyr Glu Asp Asp Val Gly Phe Gly Gly 20 25 30	
Tyr Gly Gly Tyr Gly Ala Gly Ser Asp Tyr Lys Ser Arg Gly Pro Ser	
Thr Asn Gln Ile Leu Ala Leu Ile Ala Gly Val Pro Ile Gly Gly Thr	

Leu Leu Thr Leu Ala Gly Leu Thr Leu Ala Gly Ser Val Ile Gly Leu 65 70 75 80

Leu Val Ser Ile Pro Leu Phe Leu Leu Phe Ser Pro Val Ile Val Pro

Ala Ala Leu Thr Ile Gly Leu Ala Val Thr Gly Ile Leu Ala Ser Gly 105

Leu Phe Gly Leu Thr Gly Leu Ser Ser Val Ser Trp Val Leu Asn Tyr 120

Leu Arg Gly Thr Ser Asp Thr Val Pro Glu Gln Leu Asp Tyr Ala Lys 135

Arg Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Met Lys Gly Lys Glu 160

Met Gly Gln Tyr Val Gln Asp Lys Ala His Glu Ala Arg Glu Thr Glu 165 170

Phe Met Thr Glu Thr His Glu Pro Gly Lys Ala Arg Arg Gly Ser 180 185

(2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 191 amino acids

 - (B) TYPE: amino acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Ala Asn Val Asp Arg Asp Arg Val His Val Asp Arg Thr Asp

Lys Arg Val His Gln Pro Asn Tyr Glu Asp Asp Val Gly Phe Gly Gly

Thr Gly Gly Thr Gly Ala Gly Ser Asp Tyr Lys Ser Arg Gly Pro Ser

Thr Asn Gln Ile Leu Ala Leu Ile Ala Gly Val Pro Ile Gly Gly Thr

Leu Ile Thr Leu Ala Gly Leu Thr Leu Ala Gly Ser Val Ile Gly Leu

Leu Val Ser Ile Pro Leu Phe Leu Ile Phe Ser Pro Val Ile Val Pro 85

Ala Ala Leu Thr Ile Gly Leu Ala Val Thr Gly Ile Leu Ala Ser Gly 110

Leu Phe Gly Leu Thr Gly Leu Ser Ser Val Ser Trp Val Leu Asn Tyr 115 120 125

Leu Arg Gly Thr Ser Asp Thr Val Pro Glu Gln Leu Asp Tyr Ala Lys
130 135 140

Arg Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Met Lys Gly Lys Glu 145 150 155 160

Met Gly Gln Tyr Val Gln Asp Lys Ala His Glu Ala Arg Glu Thr Glu
165 170 175

Phe Met Thr Glu Thr His Glu Pro Gly Lys Ala Arg Arg Gly Ser 180 185 190

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 191 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Ala Asn Val Asp Arg Asp Arg Val His Val Asp Arg Thr Asp 1 5 10 15

Lys Arg Val His Gln Pro Asn Tyr Glu Asp Asp Val Gly Phe Gly Gly 20 25 30

Thr Gly Gly Thr Gly Ala Gly Ser Asp Tyr Lys Ser Arg Gly Pro Ser 35 40 45

Thr Asn Gln Ile Leu Ala Leu Ile Ala Gly Val Pro Ile Gly Gly Thr 50 60

Leu Ile Thr Leu Ala Gly Leu Thr Leu Ala Gly Ser Val Ile Gly Leu 65 70 75

Leu Val Ser Ile Pro Leu Phe Leu Ile Phe Ser Pro Val Ile Val Pro 85 90 95

Ala Ala Leu Thr Ile Gly Leu Ala Val Thr Gly Ile Leu Ala Ser Gly 100 105 110

Leu Phe Gly Leu Thr Gly Leu Ser Ser Val Ser Trp Val Leu Asn Tyr 115 120 125

Leu Arg Gly Thr Ser Asp Thr Val Pro Glu Gln Leu Asp Tyr Ala Lys
130 135 140

Arg Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Met Lys Gly Lys Glu 145 150 155 160 Met Gly Gln Tyr Val Gln Asp Lys Ala His Glu Ala Arg Glu Thr Glu 165 170 175

Phe Met Thr Glu Thr His Glu Pro Gly Lys Ala Arg Arg Gly Pro 180 185 190

(2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 78 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Phe Gly Leu Thr Gly Leu Ser Ser Val Ser Trp Val Leu Gln Leu Pro
1 10 15

Pro Trp Ala Ser Asp Thr Val Pro Glu Gln Val Asp Tyr Ala Lys Arg 20 25 30

Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Met Lys Gly Lys Glu Met 35 40 45

Gly Gln Tyr Val Gln Asp Lys Ala His Glu Ala Arg Glu Thr Glu Phe 50 60

Met Thr Glu Thr His Glu Pro Gly Lys Ala Arg Arg Gly Ser 65 70 75

(2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 173 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Met Ala Asp Thr Ala Arg Gly Thr His His Asp Ile Ile Gly Arg Asp 1 10 15

Gln Tyr Pro Met Met Gly Arg Asp Arg Asp Gln Tyr Gln Met Ser Gly 20 25 30

Arg Gly Ser Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Ala Thr 35 40 45

Ala Val Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Ser Leu Thr Leu 50 60

Val Gly Thr Val Leu Ala Leu Thr Val Ala Thr Pro Leu Leu Val Leu 65 70 75 80

Phe Ser Pro Ile Leu Val Pro Ala Leu Ile Thr Val Ala Leu Leu Ile 85 90 95

Thr Gly Phe Leu Ser Ser Gly Gly Phe Gly Ile Ala Ala Ile Thr Val 100 105 110

Phe Ser Trp Ile Tyr Lys Tyr Ala Thr Gly Glu His Pro Gln Gly Ser

Asp Lys Leu Asp Ser Ala Arg Met Lys Leu Gly Ser Lys Ala Gln Asp 130 135 140

Leu Lys Asp Arg Ala Gln Tyr Tyr Gly Gln Gln His Thr Gly Gly Glu
145 150 155 160

His Asp Arg Asp Arg Thr Arg Gly Gly Gln His Thr Thr

(2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 141 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Met Ala Asp Gln Thr Arg Thr His His Glu Met Ile Ser Arg Asp Ser 1 5 10 15

Thr Gln Glu Ala His Pro Lys Ala Arg Gln Trp Val Lys Ala Ala Thr

Ala Val Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Gln Leu Thr Leu 35 40 45

Ala Gly Thr Val Ile Ala Leu Thr Val Ala Thr Pro Leu Leu Val Ile 50 60

Phe Ser Pro Val Leu Val Pro Ala Val Val Thr Val Ala Leu Ile Ile 65 70 75 80

Thr Gly Phe Leu Ala Ser Gly Gly Phe Gly Ile Ala Ala Ile Thr Ala 85 90 95

- Phe Ser Trp Leu Tyr Arg His Trp Thr Gly Ser Gly Ser Asp Lys Ile 100 105 110
- Glu Trp Ala Arg Met Lys Val.Gly Ser Arg Val Gln Asp Thr Lys Tyr
 115 120 125
- Gly Gln His Trp Ile Gly Val Gln His Gln Gln Val Ser 130 135 140

(2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 199 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Met Ala Asp Thr His Arg Val Asp Arg Thr Asp Arg His Phe Gln Phe 1 15
- Gln Ser Pro Tyr Glu Gly Gly Arg Gly Gln Gly Gln Tyr Glu Gly Asp 20 25 30
- Arg Gly Tyr Gly Gly Gly Gly Tyr Lys Ser Met Met Pro Glu Ser Gly 35
- Pro Ser Ser Thr Gln Val Leu Ser Leu Leu Ile Gly Val Pro Val Val 50 60
- Gly Ser Leu Ile Ala Leu Ala Gly Leu Leu Leu Ala Gly Ser Val Ile
 65 70 75 80
- Gly Leu Met Val Λ la Leu Pro Leu Phe Leu Ile Phe Ser Pro Val Ile 85 90 95
- Val Pro Ala Gly Leu Thr Ile Gly Leu Ala Met Thr Gly Phe Leu Ala
 100 105 110
- Ser Gly Met Phe Gly Leu Thr Gly Leu Ser Ser Ile Ser Trp Val Met 115 120 125
- Asn Tyr Leu Arg Gly Thr Ala Arg Thr Val Pro Glu Gln Leu Glu Tyr 130 135 140
- Ala Lys Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Gln Lys Gly
 145 150 155 160
- Lys Glu Met Gly Gln His Val Gln Asn Lys Ala Gln Asp Val Lys Gln 165 170 175

Tyr Asp Ile Ser Lys Pro His Asp Thr Thr Thr Lys Gly His Glu Thr 180 185 190

Gln Gly Gly Thr Thr Ala Ala 195

(2) INFORMATION FOR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 199 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
- Met Ala Asp Thr His Arg Val Asp Arg Thr Asp Arg His Phe Gln Phe 1 5 10 15
- Gln Ser Pro Tyr Glu Gly Gly Arg Gly Gln Gly Gln Tyr Glu Gly Asp
 20 25 30
- Arg Gly Tyr Gly Gly Gly Gly Tyr Lys Ser Met Met Pro Glu Ser Gly 35 40 45
- Pro Ser Ser Thr Gln Val Leu Ser Leu Leu Ile Gly Val Pro Val Val 50 60
- Gly Ser Leu Ile Ala Leu Ala Gly Leu Leu Ile Ala Gly Ser Val Ile 65 70 75 80
- Gly Leu Met Val Ala Leu Pro Leu Phe Leu Ile Phe Ser Pro Val Ile 85 90 95
- Val Pro Ala Ala Leu Thr Ile Gly Leu Ala Met Thr Gly Phe Leu Ala 100 105 110
- Ser Gly Met Phe Gly Leu Thr Gly Leu Ser Ser Ile Ser Trp Val Met
- Asn Tyr Leu Arg Gly Thr Arg Arg Thr Val Pro Glu Gln Leu Glu Tyr 130 135 140
- Ala Lys Arg Arg Met Ala Asp Ala Val Gly Tyr Ala Gly Gln Lys Gly
 150 155 160
- Lys Glu Met Gly Gln His Val Gln Asn Lys Ala Gln Asp Val Lys Gln 165 170 175
- Tyr Asp Ile Ser Lys Pro His Asp Thr Thr Thr Lys Gly His Glu Thr 180 185 190

Gln Gly Arg Thr Thr Ala Ala 195

(2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1267 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

					TGCAGACCAA	60
GAGAAAGTG?	A GAGAGAGTTG	TCCTCTCGTT	ATCAÄGTAAC	AGTAGACCAC	CACTAAACCG	120
	ТАТАЛТСАЛАЛ					180
TCCATGGACT	GCCTACCCGA	ATTGATTGAT	TCGACTAGTT	TTTCTTCTTC	TTTGATTAAG	240
ACCTCCGTAA	GAAAAATGGT	ACTACTAAAG	CCACTCGCTA	ССААА́АСТАА	ACCATTCCAG	300
					AACTAATTAA	360
GAACTATAAC	CATTAACCGT	АААААТАААТ	TTACTACAGT	AAAAAATTAT	ACTAATTTCA	420
	AATTTCAGCT					480
	ATCCTTATTT					540
	AATTCTATAT					600
	ATACTCTTGT			•		660
TCATCCTACA	TTCATACCTA	AGCTAGCAAA	GCAAACTACT	AAAAGGGTCG	TCAACGCAAG	720
TTATTTGCTA	GTTGGTGCAT	ACTACACACG	GCTACGGCAA	CATTAAGTAA	CACATTAAGA	780
	TAATGTAGTA					840
GGTACAGGTA	GATGAAAAAT	ATTTGGTTAG	CGGGTTGAGA	TTAAGCGGAT	ATAGGAGGCA	900
TATATACAGC	TGTGAGAAGA	AGAGGGATAA	АТЛСААЛАА G	GGAAGGATGT	TTTTGCCGAC	960
NGAGAAAGGT	AGATTAAGTA	GGCATCGAGA	GGAGAGCAAT	TGTAAAATGG	ATGATTTGTT	1020
TGGTTTTGTA	CGGTGGAGAG	AAGAACGAAA	AGATGATCAG	GTAAAAAATG	AAACTTGGAA	1080
ATCATGCAAA	GCCACACCTC	TCCCTTCAAC	ACAGTCTTAC	GTGTCGTCTT	CTCTTCACTC	1140
САТАТСТССТ	TTTTATTACC	AAGAAATATA	TGTCAATCCC	ATTTATATGT	ACGTTCTCTT	1200

AGACTTA	TCT CTATATACCC CCTTTTAATT TGTGTGCTCT TAGCCTTTAC TCTATAGTTT	1260
TAGATAG		1260
		1267
(2) INF	ORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
GTAATAC	GAC TCACTATAGG GC	22
(2) INF(ORMATION FOR SEQ ID NO:14:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA (genomic)	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:14:	
GGGATCC	TA TACTAAAACT ATAGAGTAAA GG	32
(2) INFO	RMATION FOR SEQ ID NO:15:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 8 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	MOLECULE TYPE: protein	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:15:	
Trp 1	Ile Gly His Asp Ala Gly His 5	

- (2) INFORMATION FOR SEQ ID NO:16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Asn Val Gly His Asp Ala Asn His 1

- (2) INFORMATION FOR SEQ ID NO:17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Val Leu Gly His Asp Cys Gly His 1

- (2) INFORMATION FOR SEQ ID NO:18:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Val Ile Ala His Glu Cys Gly His

- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Val Ile Gly His Asp Cys Ala His 1

- (2) INFORMATION FOR SEQ ID NO:20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Val Val Gly His Asp Cys Gly His 1

- (2) INFORMATION FOR SEQ ID NO:21:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

His Asn Ala His His
1 5

- (2) INFORMATION FOR SEQ ID NO:22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

His Asn Tyr Leu His His 1

- (2) INFORMATION FOR SEQ ID NO:23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

His Arg Thr His His

- (2) INFORMATION FOR SEQ ID NO:24:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

His Arg, Arg His His

- (2) INFORMATION FOR SEQ ID NO:25:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

His Asp Arg His His

- (2) INFORMATION FOR SEQ ID NO:26:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

His Asp Gln His His

- (2) INFORMATION FOR SEQ ID NO:27:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

His Asp His His His

- (2) INFORMATION FOR SEQ ID NO:28:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids

 - (B) TYPE: amino acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

His Asn His His His

- (2) INFORMATION FOR SEQ ID NO:29:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

Phe Gln Ile Glu His His

- (2) INFORMATION FOR SEQ ID NO:30:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

His Gln Val Thr His His 1

- (2) INFORMATION FOR SEQ ID NO:31:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

His Val Ile His His 1 5

- (2) INFORMATION FOR SEQ ID NO:32:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

His Val Ala His His 1 5

- (2) INFORMATION FOR SEQ ID NO:33:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE	TYPE:	protein
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

His Ile Pro His His 5

(2) INFORMATION FOR SEQ ID NO:34:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

His Val Pro His His 1

(2) INFORMATION FOR SEQ ID NO:35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1941 base pairs

 - (B), TYPE: nucleic acid (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

GAGCTCGATC	ACACAAAGAA	AACGTCAAAT	GGATCATACT	GGGCCCATTT	TGCAGACCAA	60
GAGAAAGTGA	GAGAGAGTTG	TCCTCTCGTT	ATCAAGTAAC	AGTAGACCAC	CACTAAACCG	120
CCAATAGCTT	ATAATCAAAA	TAGAAAGGTC	TAATAACAGA	AACAAATGAA	AAAGCCTTGT	180
TCCATGGACT	GCCTACCCGA	ATTGATTGAT	TCGACTAGTT	TTTCTTCTTC	TTTGATTAAG	240
ACCTCCGTAA	GAAAAATGGT	ACTACTAAAG	CCACTCGCTA	ССААААСТАА	ACCATTCCAG	300
ACTGTAACTG	GACCAATATT	TCTAAACTGT	AACCAGATCT	САААСАТАТА	ААСТААТТАА	360
GAACTATAAC	CATTAACCGT	ААЛААТАААТ	TTACTACAGT	AAAAAATTAT	ACTAATTTCA	420
GCTATGATGG	AATTTCAGCT	CTTAAGAGTT	GTGGAAATCA	AGTAAACCTA	AAATCCTAAT	480
AATATTCTTC	ATCCTTATTT	TTGTTTCACA	TGCATGCTGT	CCAATCTGTT	ATTAGCATTT	540
GAAAGCCTAA	AATTCTATAT	ACAGTACAAT	AAATCTAATT	AATTTTCATT	АСТААТАЛАА	600

TGCTTCATAT A	TACTCTTGT	ATTTATAAAT	CATCCGTTAT	CGTTACTATA	CCTTTATACA	660
TCATCCTACA T	TCATACCTA	AGCTAGCAAA	GCAAACTACT	AAAAGGGTCG	TCAACGCAAG	720
TTATTTGCTA G	TTGGTGCAT	ACTACACACG	GCTACGGCAA	CATTAAGTAA	CACATTAAGA	780
GGTGTTTTCT T	'AATGTAGTA	TGGTAATTAT	ATTTATTTCA	AAACTTGGAT	TAGATATAAA	840
GGTACAGGTA G	ATGAAAAAT	ATTTGGTTAG	CGGGTTGAGA	TTAAGCGGAT	ATAGGAGGCA	900
TATATACAGC T	GTGAGAAGA	AGAGGGATAA	,ATACAAAAAG	GGAAGGATGT	TTTTGCCGAC	960
AGAGAAAGGT A	GATTAAGTA	GGCATCGAGA	GGAGAGCAAT	TGTAAAATGG	ATGATTTGTT	1020
TGGTTTTGTA C	GGTGGAGAG	AAGAACGAAA	AGATGATCAG	GTAAAAAATG	AAACTTGGAA	1080
ATCATGCAAA G	CCACACCTC	TCCCTTCAAC	ACAGTCTTAC	GTGTCGTCTT	CTCTTCACTC	1140
CATATCTCCT T	TTTATTACC	ААСАААТАТА	TGTCAATCCC	ATTTATATGT	ACGTTCTCTT	1200
AGACTTATCT C	ТАТАТАССС	CCTTTTAATT	TGTGTGCTCT	TAGCCTTTAC	TCTATAGTTT	1260
TAGATAGACA TO	GGCGAATGT	GGATCGTGAT	CGGCGTGTGC	ATGTAGACCG	TACTGACAAA	1320
CGTGTTCATC AC	GCCAAACTA	CGAAGATGAT	GTCGGTTTTG	GTGGCTATGG	CGGTTATGGT	1380
GCTGGTTCTG AT	ITATAAGAG	TCGCGGCCCC	TCCACTAACC	AAGTATTTTT	GTGGTCTCTT	1440
TAGTTTTTCT TO	GTGTTTTCC	TATGATCACG	CTCTCCAAAC	TATTTGAAGA	TTTTCTGTAA	1500
ATTCATTTTA A	ACAGAAAGA	ТАААТАААТ	AGTGAAGAAC	CATAGGAATC	GTACGTTACG	1560
TTAATTATTT CO	CTTTTAGTT	CTTAAGTCCT	AATTAGGATT	ССТТТАЛААС	TTGCAACAAT	1620
CTAATTGTTC AC	CAAAATGAG	TAAAGTTTGA	AACAGATTTT	TATACACCAC	TTGCATATGT	1680
TTATCATGGT G	ATGCATGCT	TGTTAGATAA	ACTCGATATA	ATCAATACAT	GCAGATCTTG	1740
GCACTTATAG CA	AGGAGTCCA ,	TTGGTGGCAC	ACTGCTAACC	CTAGCTGGAC	TCACTCTAGC	1800
CGGTTCGGTG AT						1860
AGTCCCGGCG GC						1920
TGGGTTGACG GG					-	1941

What is claimed is:

- 1. An isolated nucleic acid encoding an oleosin 5' regulatory region which directs seed-specific expression selected from the groups consisting of the nucleotide sequence set forth in SEQ ID NO:12, the nucleotide sequence set forth in SEQ ID NO:12 having an insertion, deletion, or substitution of one or more nucleotides, or a contiguous fragment of the nucleotide sequence set forth in SEQ ID NO:12.
- 2. An expression cassette which comprises the oleosin 5' regulatory region of Claim 1 operably linked to at least one of a nucleic acid encoding a heterologous gene or a nucleic acid encoding a sequence complementary to a native plant gene.
- 3. The expression cassette of Claim 2 wherein the heterologous gene is at least one of a fatty acid synthesis gene or a lipid metabolism gene.
- 4. The expression cassette of Claim 3 wherein the heterologous gene is selected from the group consisting of an acetyl-coA carboxylase gene, a ketoacyl synthase gene, a malonyl transacylase gene, a lipid desaturase gene, an acyl carrier protein (ACP) gene, a thioesterase gene, an acetyl transacylase gene, or an elongase gene.
- 5. The expression cassette of Claim 4 wherein the lipid desaturase gene is selected from the group consisting of a $\Delta 6$ -desaturase gene, a $\Delta 12$ -desaturase gene, and a $\Delta 15$ -desaturase gene.
- 6. An expression vector which comprises the expression cassette of any one of Claims 2-5.

- 7. A cell comprising the expression cassette of any one of Claims 2-5.
- 8. A cell comprising the expression vector of Claim 6.
- 9. The cell of Claim 7 wherein said cell is a bacterial cell or a plant cell.
- 10. The cell of Claim 8 wherein said cell is a bacterial cell or a plant cell.
- 11. A transgenic plant comprising the expression cassette of any one of Claims 2-5.
- 12. A transgenic plant comprising the expression vector of Claim 6.
- 13. A plant which has been regenerated from the plant cell of Claim 9.
- 14. A plant which has been regenerated from the plant cell of Claim 10.
- 15. The plant of Claim 12 or 13 wherein said plant is at least one of a sunflower, soybean, maize, cotton, tobacco, peanut, oil seed rape or Arabidopisis plant.
 - 16. Progeny of the plant of Claim 11 or 12.
 - 17. Seed from the plant of Claim 11 or 12.
- 18. A method of producing a plant with increased levels of a product of a fatty acid synthesis gene or a lipid metabolism gene which comprises:
- (a) transforming a plant cell with an expression vector comprising the isolated nucleic acid of Claim 1 operably linked to at least one of an

isolated nucleic acid coding for a fatty acid synthesis gene or a lipid metabolism gene; and

- (b) regenerating a plant with increased levels of the product of said fatty acid synthesis or said lipid metabolism gene from said plant cell.
- 19. A method of producing a plant with increased levels of gamma linolenic acid (GLA) content which comprises:
- (a) transforming a plant cell with an expression vector comprising the isolated nucleic acid of Claim 1 operably linked to a $\Delta 6$ -desaturase gene; and
- (b) regenerating a plant with increased levels of GLA from said plant cell.
- 20. The method of Claim 19 wherein said $\Delta 6$ -desaturase gene is at least one of a cyanobacterial $\Delta 6$ -desaturase gene or a Borage $\Delta 6$ -desaturase gene.
- 21. The method of any one of Claims 18-20 wherein said plant is a sunflower, soybean, maize, tobacco, cotton, peanut, oil seed rape or Arabidopsis plant.
- 22. The method of Claim 18 wherein said fatty acid synthesis gene or said lipid metabolism gene is at least one of a lipid desaturase, an acyl carrier protein (ACP) gene, a thioesterase gene an elongase gene, an acetyl transacylase gene, an acetyl-coA carboxylase gene, a ketoacyl synthase gene, or a malonyl transacylase gene.
- 23. A method of inducing production of at least one of gamma linolenic acid (GLA) or

octadecatetraeonic acid (OTA) in a plant deficient or lacking in GLA which comprises transforming said plant with an expression vector comprising an the isolated nucleic acid of Claim 1 operably linked to a $\Delta6$ -desaturase gene and regenerating a plant with increased levels of at least one of GLA or OTA.

- 24. A method of decreasing production of a fatty acid synthesis or lipid metabolism gene in a plant which comprises:
- (a) transforming a plant cell with an expression vector comprising the isolated nucleic acid of Claim 1 operably linked to a nucleic acid sequence complementary to a fatty acid synthesis or lipid metabolism gene; and
- (b) regenerating a plant with decreased production of said fatty acid synthesis or said lipid metabolism gene.
- 25. A method of cosuppressing a native fatty acid synthesis or lipid metabolism gene in a plant which comprises:
- (a) transforming a cell of the plant with an expression vector comprising the isolated nucleic acid of Claim 1 operably linked to a nucleic acid sequence encoding a fatty acid synthesis or lipid metabolism gene native to the plant; and
- (b) regenerating a plant with decreased production of said fatty acid synthesis or said lipid metabolism gene.

ata tot god tac cot coe ass gag agt agt cat tit toa toa atg got got cas ate a HAAOIK as a tac att acc tre det das che as acc car dat as ccc dos dat cre top atc tre tre R γ I T S D S L R II II D R P G D L W I S 122 102 ccc ttg aag agt ctt gct ggt caa gag gta act gat gce ttt gtt gcs tte cat cct gcc P L K S L A G Q & V T D A F V A F H P A 242 tet aca top and nat ctt dat and til the act gop tot tot ett ann gat tac tel git 103 KHLOKFF S G V S K D Y R K L V F E F S K H O L Y 362 gas as an ggt cat stt atg tit gen act tig tgc tit ata gen atg cig tit get atg D K K G H I H F λ T L C F I λ M L F λ M 422 aut git tot gog get etg tet tot gog got get etg gen cat etg tet tet gog tot etg S V Y G V L F C G G V L V H L F S G C L 442 542 Plotet gat tea agg ett aat aag itt atg ggl att itt get gea aat igt ett tea gga V S D S R I. H K F H G 1 F A A H C L S G 407 ate agt att got tog tog eas tog eac cat eat goe cat cac att goe tot eat ege ctt I .S I G W W K W W II H A II II I A C H S L 662 gas tat gas cot gat tta cas tat ata oca tts ott gtt gtg tot toe aag ttt ttt ggt E Y D P D L O Y 1 P F L V V S S K F F G tes etc acc tet cat tre tat gag ass agg tre act tre gac tet tra tea aga tre tre S L T S H F Y E R R L T F D S L S R F F 787 gta agt tat cas cat top aca tit tac cot att atg tot got got agg ete aat atg tat 2 A O U A TFYPIH 843 V O S f. I H f. L T K R N V ges can tet etc ata std ttd ttd acc and aga nat gtd tee tat con get con gas etc S Y tto gra tgc cta gtg ttc trg att tgg tag ccg ttg ctt gtt tct tgc ttg cct aat tgg L G C L V F S 1 M Y P 1, L V S C L P $\rm H$ M 362 G F R I H F V I A S L S V T G H Q O V Q 1022 tto too too sac ose the tet tea sot oft tat out one say cet ass god sat sat too F . S . S . V . V . G . F . F . G . H . W1082 tit gag ass cas acg gat ggg aca cit gam att tot tot not not tog atg gat tog tit EKOTOGTLDISCIP cat ggt gga tig can tic can att gag cat cat tig til ecc ang atg cet aga tgc asc G L O F O L P II II L F P K H P R C H 1207 ctt agg aam ate teg eee tae gtg ate gag tta tge aag ama cat mat tig eet tae amt KISPYV I E L C K K H H L P Y 1262 tat gos tot the toe say one sat was also see one one are the sag are the sage are sages and sages are sages are sages and sages are sages 1322 Cad jet agg gat ata acc eag ceg etc ecg aag aat tty gta tgg gaa get ett eac act Q A R D I T K P L P K N L V W E A L II T eat ggt tam ast tac ect tag tto stg tam too ttt gag att stg tat etc eta tgt ttg 1442 tor cet gen ten get cen cet get one gen are one ect tor cet ten egg ter att en 1502 tot tet the are tot til age got til got til et tot ett att get gee tee goe git per tat the costen ern too too sta tot gat att ten gas tot act ten too can tot 1622 get the age toa age tea tot gea ett era tag act tog the aga tog tha tot cat oth att L

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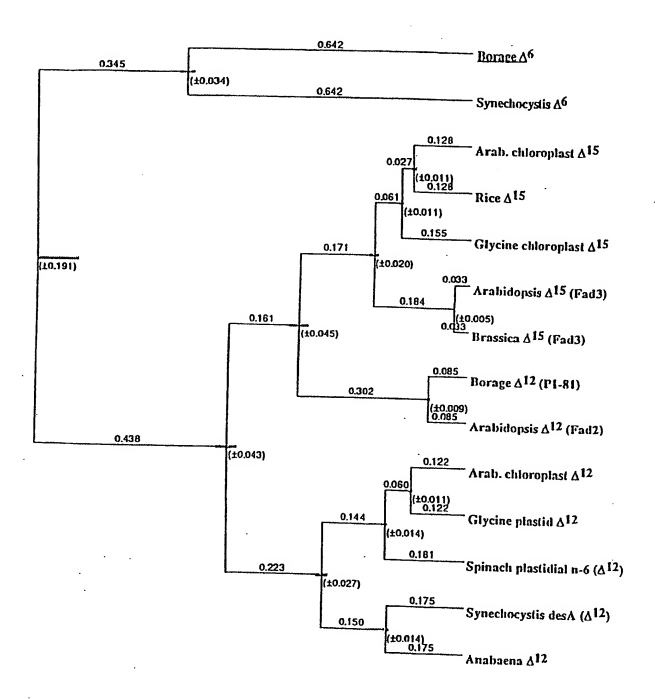


FIGURE 2

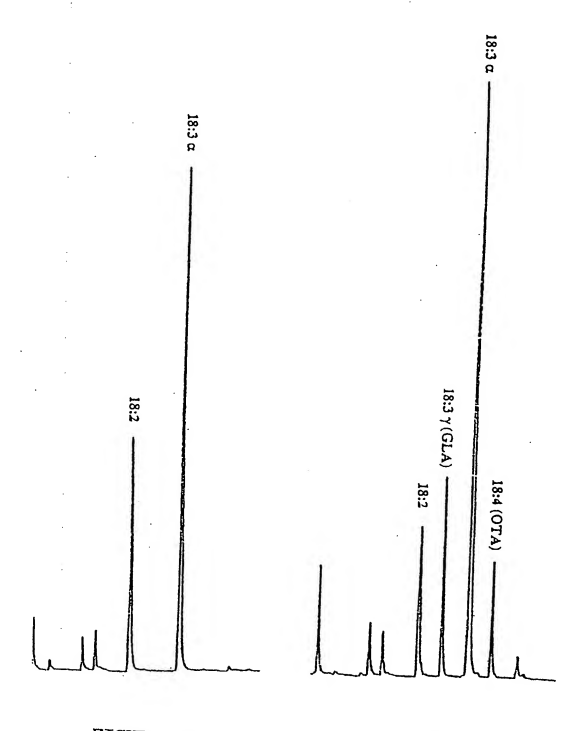


FIGURE 3A

FIGURE 3B

FIGURE 4

1 tta god ttt act cta tag ttt tag ata gad atg gog aat gtg gat ogt gat ogg ogt gtg H A N V D R D R R V 61 cat gta gao cgt act gac ama cgt gtt cat cag coa aac tao gaa gat gat gto ggt ttt 11 H V D R T D K R V H Q P N Y B D D V G F 121 ggt ggo tat ggc ggt tat ggt gct ggt tct gat tat aag agt cgc ggc ccc tcc act aac BIOG Y G G Y G A G S D Y K S R G P S T H 101 can ato ttg gos oft ats gos ggs gtt coo att ggt ggc acs etg ots acc ets get ggs 51 Q I L A L I A G V P I G G T L L T L A G 241 etc act eta gee ggt teg gtg ate gge ttg eta gte tee ata ecc etc tte etc etc tte 71 L T L A G S V I G L L V S I P L F L L F 301 agt ccg gtg ata gtc ccg gcg gct ctc act att ggg ctt gct gtg acg gga atc ttg gct VIVPAALTIGLAVTCILA 91 S P 361 tot ggt tig tit ggg tig acg ggt cig ago tog gto tog tgg gto oto aac tao oto egt HISGLFCLTCLSSVSWVLRYLR 421 ggg acg agt gat aca gtg cca gag caa ttg gac tac gct aan cgg cgt atg gct gat gcg. 131 G T S D T V P E Q L D Y A K R R H A D A 481 gta ggc tat gct ggt atg aag gga aaa gag_atg ggt cag tat gtg caa gat aag gct cat 151 V G Y A G H K G K E H G Q Y V Q D K A H 541 gag got ogt gag act gag tto atg act gag acc cat gag cog ggt aag goc agg aga ggc 171 EARETEFHTETBEPGKARRG 601 toa taa got aat ata aat tgo ggg agt cag ttg gaa acg oga taa atg tag ttt tac ttt 191 Ş · 661 tat gto com gtt tot tto oto tit tam gam tat ott tgt otm tat atg tgt tog tto gtt 721 ttg tct tgt cca aat aas aat cct tgt tag tga aat aag aas tga aat aas tat gtt ttc 781 ttt ttt gag ata acc aga sat ctc ata cta ttt tct ana ana ana ana ana ana

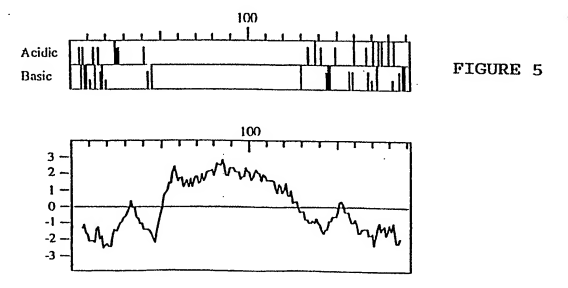


FIGURE 6

ALS21	HNNVDRDRRV	BVDRTDKRVB	-QPNYEDDVG	P-GGY	GGYGAGSDYK	43
X91918	HANVDRDRRV	BVDRTDKRVB	-QPNYEDDVG	F-GGY	GGYBAGSDYK	43
229859						
x62352	HIDTARG	OFFICERT-	OXPHINER DEED	Duesangro	SDT-	37
Atol3	MADORR					-21
x91956	MADTHR					45
L40954	инотня	-VDRTDR8FQ	FQSPYEGGRG	QGQYEGDRGY	GGGGYRSHMP	45
AtS21	SRGPSTNOIL			भवन्यावाप्यक	Teletresea	93
X91918	SRGPSTNIL	ALIAGUPIGG	TLIPRAGLTI	Machine 1	Denernesen	93
229859			++++	++++++	++-+-+	
X62352	akandra			ANTHALTHAN .		84
Atol3	BRARCES	RAATAVITAG	arimiseri	AVILIATIVE	161 CA1626	68
X91956	ESGPSSICVL			AVMIDIVEDA		95
L40954	ESGPSSIOVL	2PridAbana	arrivarrii	MANATHATA	Tellemiesen	95
AES21	IDIMIAGAN	JE Alterty	FGLTGLSSVS	WILLIAMS	DTVPEOLDYA	143
X91918				Wyleniens		143
229859			FGLTGLSSVS	WALDERS SEE	DTVPEOVDYA	30
X62352	IVPACTIVAL	r. irdelusses				134
Atol3	INPASSIVAL			WLYSERTGS-		115
X91956	IVPAGINICI			WHHYLRGTA		145
L40954	DINTANAL				RTVPEQLEYA	145
AtS21	WDDMADAUGV	ACHERRHOO	VVODKAHEAR	ETEF	MTĖTHEPGKA	187
X91918					MTETHEPGKA	187
229859	KRRHADAVGY	AGHRERENGO	YVODKAHEAR	ETEF	MTETHEPGKA	74
X62352					23	173
Atol3					83	141
X91956	KDBMVUVACA	PCCKERENCO	RVOOKAOOK	OYDISKPHOT	TTKGHETQGB	195
L40954		7 1	7.71		TTKGBETQGA	195
AtS21	RRG# = 254	164/Atol2		•		191
X91918	RRGP	10.1/11077			•	191
Z29859	RRGS					78
X62352	-4/00					173
Atol3						141
X91956	TTAA					199
L40954	TTAA					199
240724	- 1101					200

FIGURE 7

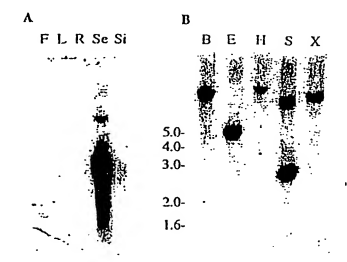


FIGURE 8B

FIGURE 8A

FIGURE 9

٨ 1 81 161 ACCTCCGTAAGAAAATGGTACTAAAGCCACTCGCTACCAAAACTAAACCATTCCAGACTGTAACTGGACCAATATT 241 321 AAAAAATTATACTAATTCAGCTATGATGGAATTTCAGCTCTTAAGAGTTGTGGAAATC.AGTAAAACCTAAAATCCTAAT 401 AATATTCTTCATCCTTATTTTTGTTTCACATCCATGCTGTCCAATCTGTTATTAGCATTTGAAAGCCTAAAATTCTATAT 4B1 ACAGTACAATAAATCTAATTAATTTCATTACTAATAAAATCCTTCATATATACTCTTGTA<u>TTTATAAATCATCC</u>GTTAT 561 CGTTACTATACC<u>TTTATACATCATCC</u>TACATTCATACCTAAGCTAGCAAACTACTAAAAGGGTCGTCAACGCAAG 641 TTATTTGCTAGTTGGTGCATACTACACACGGCTACGGCAACATTAAGTAACACATTAAGAGGTGTTTTCTTAATGTAGTA 721 TOGTARTTATATTTCAAAACTTGGATTAGATATAAAGGTACAGGTAGATGAAAAATATTTGGTTAGGGGGTTGAGA 801 TTAAGCGGATATAGGAGGCATATATACAGCTGTGAGAAGAGGGGATAAATACAAAAAGGGAAGGATGTTTTTGCCGAC 881 961 1041 AAGAACGAAAAGATGATCAGGTAAAAAATGAAACTTGGAAATCATGCAAAGCCACACCTCTCCCTTCAACACAGTCTTAC 1121 <u>GŢĢŢC</u>GTCTTCTCTCACTCCATATCTCCTTTTTATTACCAAGAAA!BAWATGTCAATCCCATTTA<u>TA</u>TGTACGTTCTCTT 1201 AGACTTATCTCTATATACCCCCCTTTTAATTTGTGTGCTCttagCctttactctatagttttagatagacatygcgaatgt 1281 ggatcgtgatcggcgtgtgcatgtagaccgtactgacaaacgtgttcatcagccaaactacgaagatgatgtcggttttg 1361 gtggctatggcggttatggtgctggttctgattataagagtcgcggcccctccactaaccaaGTATTTTTGTCGTCTCTT 1441 TAGTTTTCTTGTGTTTTCCTATGATCACGCTCTCCAAACTATTTGAAGATTTTCTGTAAATTCATTTTAAACAGAAAGA 1521 TAAATAAAATAGTGAAGAACCATAGGAATCGTACGTTACGTTAATTATTTCCTTTTAGTTCTTAAGTCCTAATTAGGATT 1601 CCTTTANAAGTTGCAACAATCTAATTGTTCACAAAATGAGTAAAGTTTGAAACAGATTTTATACACCACTTGCATATGT 1681 TTATCATGGTGATGCATGCTTGTTAGATAAACTCGATATAATCAATACATGCAGatcttggcacttatagcaggagttcc 1761 cattggtggcacactgctaaccctagctggactcactctagccggttcggtgatcggcttgctagtctccatacccctct 1821 | tcctcctcttcagtccggtgatagtcccggcggctctcactattgggcttgctgtgacgggaatcttggcttctggtttg 1901 tttgggttgacgggtctgagctc

B

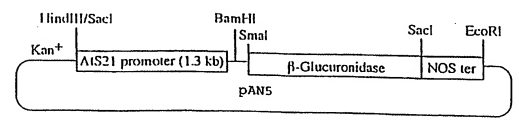


FIGURE 10

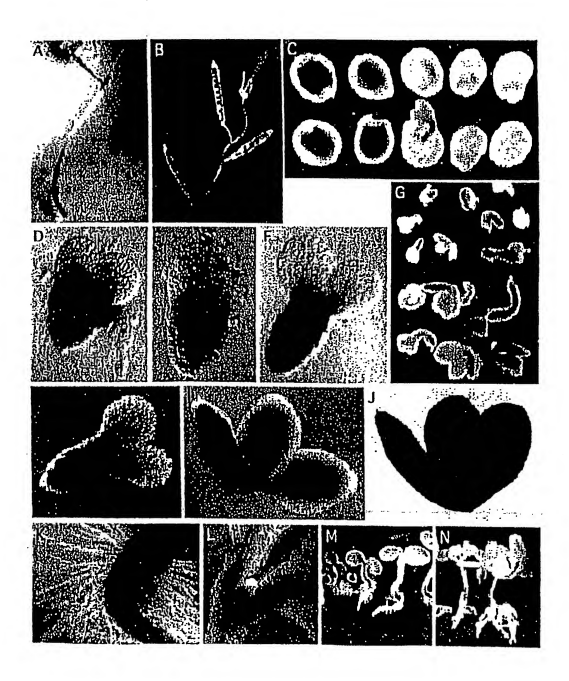


FIGURE 11A-11N

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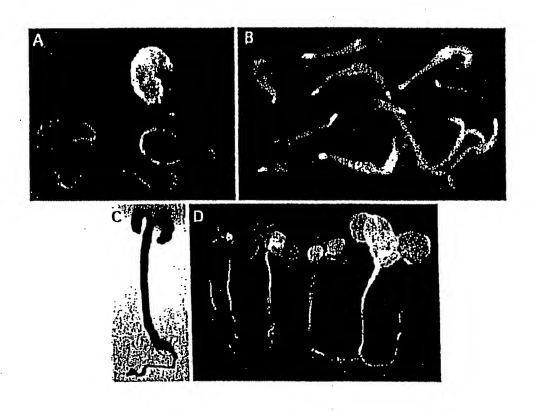


FIGURE 12A-D

FIGURE 13A

12345678 12345678 12345678



FIGURE 13B



FIGURE 13C

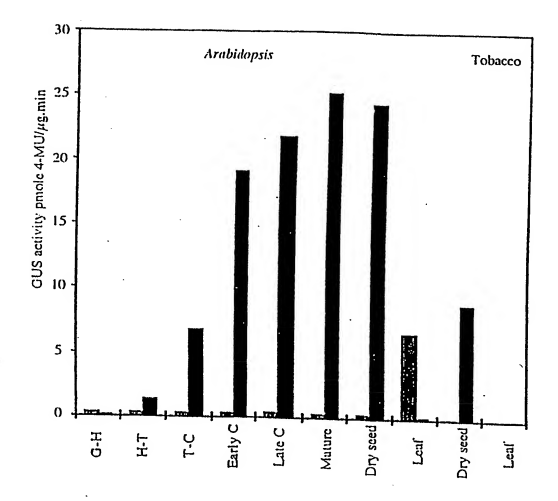
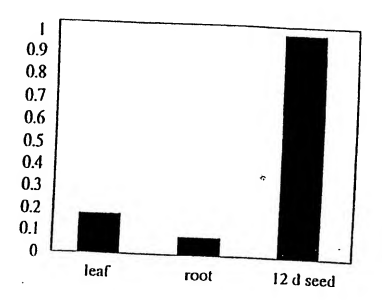


FIGURE 14

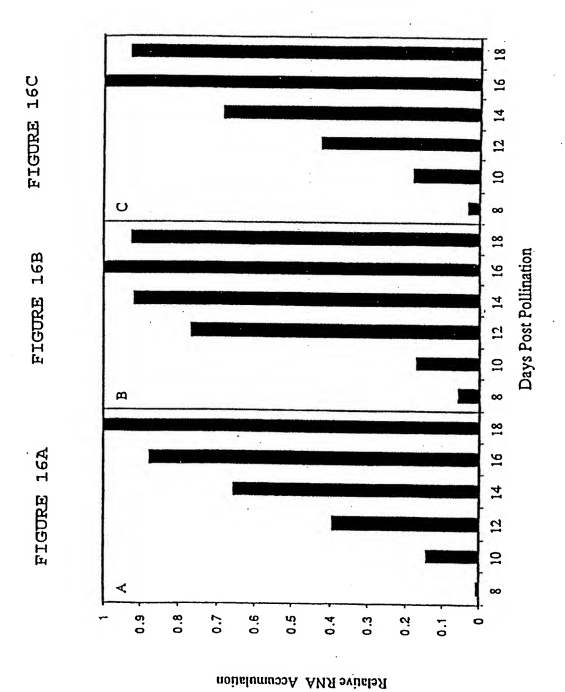
FIGURE 15B



Borage tissue



FIGURE 15A



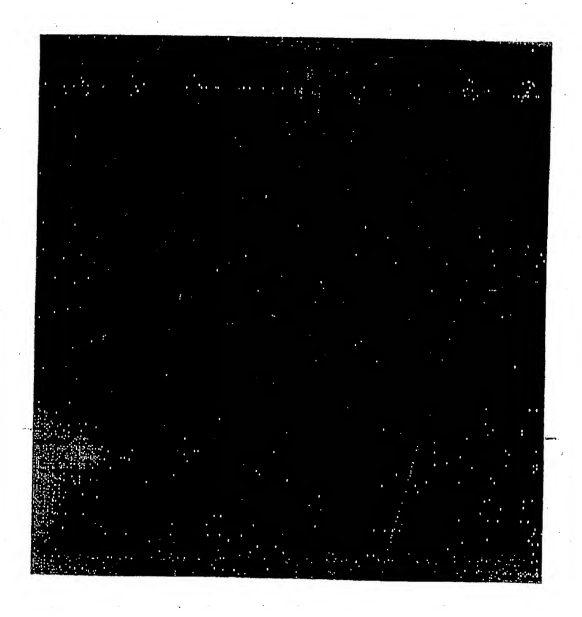
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FIGURE 17



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A. CLASS	SIFICATION OF SUBJECT MATTER			PCT/US 9	98/07179	
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According	to International Patent Classification (IPC) or to both na	ational classification an	dIPC			
B. FIELDS	SEARCHED					
IPC 6	ocumentation searched (classification system follower C12N A01H	d by classification symb	ools)	······································		
Documenta	ation searched other than minimum documentation to the	e extent that such doc	uments are include	ed in the fields :	Searched	
Electronic o	lata base consulted during the international search (ne	ame of data base and,	where practical, se	earch terms use		
	ENTS CONSIDERED TO BE RELEVANT			······································		
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	September 1998	·	21/09/1998	1		
and the	ling address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Author	rized officer			
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016 Maddox, A						

inte Jonal Application No PCT/US 98/07179

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